

POLYPEPTIDES INVOLVED IN THE BIOSYNTHESIS OF
STREPTOGRAMINS, NUCLEOTIDE SEQUENCES CODING FOR THESE
POLYPEPTIDES AND THEIR USE

The present invention relates to novel
5 polypeptides involved in the biosynthesis of
streptogramins, and also comprises the isolation and
identification of genes for the biosynthesis of the A
and B components of streptogramins, the expression of
these genes with the object of increasing the levels of
10 production and their use for the construction of
blocked mutants capable of leading to the synthesis of
novel antibiotics or to derived forms of
streptogramins.

Streptogramins form a homogeneous group of
15 antibiotics, consisting of a combination of two types
of molecules which are chemically different; on the one
hand polyunsaturated macrolactones (A-group components,
two examples of structures of which are presented in
Figure 1), and on the other hand depsipeptides (B-group
20 components, three examples of the structure of which
are presented in Figure 2). This group comprises many
antibiotics (see Table 1 and Figure 3), which are known
by different names in accordance with their origin,
including pristinamycins, mikamycins and virginiamycins.
25 (for a review, see Cocito 1979, 1983).

The A and B components have a synergistic
antibacterial activity which can reach 100 times that
of the separate components and which, in contrast to

that of each component, is bactericidal (Cocito 1979).

This activity is more especially effective against

Gram-positive bacteria such as staphylococci and

streptococci (Cocito 1979, Videau 1982). The A and B

5 components inhibit protein synthesis by binding to the

50S subunit of the ribosome (Cocito 1979; for a review,

see Di Giambattista et al. 1989).

Streptogramins are chiefly produced by

actinomycetes, including many streptomycetes, presented

10 in Table 1. In addition, streptogramins are also

synthesized by eukaryotes such as Micromonospora which

synthesizes vernamycins. Actinomycetes constitute a

very important group of microorganisms on account of

the large amount of secondary metabolites they produce,

15 including many antibiotics (beta-lactams,

tetracyclines, macrolides, aminoglycosides,

polyacetates and the like), herbicides, anticancer

agents, antifungal agents, immunomodulators and enzyme

inhibitors. Many biosynthesis pathways relating to

20 antibiotics belonging to miscellaneous classes as well

as other secondary metabolites such as pigments (for a

review, Chater 1990) have already been studied at the

present time in actinomycetes. An important aspect of

this group of bacteria is that the genes involved in

25 the same biosynthesis pathway, structural genes and

also resistance gene(s) and regulatory gene(s), are

grouped together physically on the chromosome,

constituting clusters which can reach more than 100 kb

(Hopwood et al. 1986a, Hopwood et al. 1986b, Hallam et al. 1988, Anzai et al. 1987, Ohnuki et al. 1985). To date, no example has been found to contradict this observation. Such a structural organization is of great 5 interest in the development of strategies for cloning biosynthesis genes. In effect, it is possible, starting from a single gene previously cloned by various techniques, a biosynthesis, resistance or regulatory gene, to walk along the chromosome and thus to isolate 10 the set of genes of the biosynthesis cluster.

Our knowledge of the biosynthesis pathways of each of the components of streptogramins is still very incomplete, but the origin of the different parts of each molecule has been identified by radioactive 15 labelling (Kingston et al. 1983). Thus, the A-type components are made up of two regions originating from the condensation of acetates and several amino acids such as serine and glycine, for example. As regards the B-type components, studies have shown that all the 20 amino acids present in the peptide chain are derived from natural amino acids (Hook and Vining 1973). However, no polypeptide involved in these pathways has, to date, been purified in sufficient amounts to permit its molecular characterization, and no biosynthesis 25 gene has been described. In the process of biosynthesis of the B-type components, two parts may be distinguished:

- 1) Synthesis of the precursors, or of their

analogues, of the macrocycle: 3-hydroxypicolinic acid, L-2-aminobutyric acid, p-dimethylamino-L-phenylalanine, 4-oxo-L-pipeolic acid, L-phenylglycine.

2) Formation of the macrocycle from the 5 precursors mentioned above, L-threonine and L-proline, or their analogues, with possible modification of these precursors or peptide N-methylation.

To date, only the probable metabolic origin 10 of the precursors of the macrocycle of the B-type components has been determined by studies using labelled isotopes (Reed et al., 1986, Molinero et al., 1989, Reed et al., 1989).

The present invention results from the 15 purification of polypeptides participating in the biosynthesis of streptogramins, as well as from the cloning of genes whose product participates in the biosynthesis of streptogramins. The term biosynthesis of streptogramins is understood to comprise the regulatory genes and the genes conferring resistance on 20 the producing microorganisms. Thus, the present invention makes it possible to increase the levels of production of these metabolites by means of recombinant DNA techniques. Another benefit of the present invention lies in the possibility, by construction of 25 mutants blocked in the different steps of this biosynthesis, of producing synthesis intermediates for each of the two components. These intermediates may serve as substrates for further modification for

chemical, biochemical, enzymatic or microbiological means. Similarly, isolation of the biosynthesis genes makes it possible, by gene transfer between producing strains, to manufacture hybrid antibiotics having pharmacologically advantageous properties (Hopwood *et al.*, 1985a, Hopwood *et al.*, 1985b, Hutchinson *et al.* 1989). Another benefit of the present invention lies in the fact that it provides a better knowledge of the biosynthesis pathways of the metabolites classed as streptogramins. In effect, the invention enables bacterial or fungal strains to be constructed in which one or more proteins participating in the biosynthesis of streptogramins is/are expressed under the control of suitable expression signals. Such strains may then be used to carry out bioconversions. These bioconversions may be carried out either using whole cells, or using acellular extracts of the said cells. These bioconversions may enable a streptogramin to be converted to a derived form with an enzyme of a biosynthesis pathway. For example, pristinamycin IIB may be converted in this manner to pristinamycin IIA. The same reasoning may be applied to any biosynthesis intermediate.

A first subject of the invention hence relates to a nucleotide sequence coding for a polypeptide involved in the biosynthesis of streptogramins.

More especially, several genes whose product

participates in the biosynthesis of streptogramins have been isolated from Streptomyces pristinaespiralis.

Since the streptogramins produced by this strain are more commonly designated by the term pristinamycins.

5 (see Table 1), in what follows, reference will be made in some cases to genes for the biosynthesis of pristinamycins. However, it is clear that the results obtained apply to all the streptogramins.

Pristinamycins I and II correspond, respectively, to 10 the B and A components of streptogramins. Molecules of the pristinamycin II family and of the pristinamycin I family hence designate in what follows the A and B components of streptogramins, respectively.

The present invention describes in particular 15 the isolation and characterization of the snaA, snaB, snaC, snaD, papA, papM, samS, snbA, snbC, snbD, snbE and snbR genes. These genes were isolated from a library of genomic DNA of S.pristinaespiralis. This library was obtained by partial digestion of genomic 20 DNA S.pristinaespiralis with the restriction enzyme Sau3A. Large DNA fragments, from 40 to 50 kb on average, were cloned into cosmid pHC79 (Hohn, B., and Collins, J.F., 1980). After in vitro encapsidation, E.coli strains HB101 (Boyer et Roulland-Dussoix, 1969) 25 and DH1 (Low, 1968) were transfected. The DNA library of S.pristinaespiralis thus occurs in two different strains of E.coli.

The snaA, snaB and samS (initially designated

SnaC genes are present on cosmid pIBV1 (Figure 4). The product of the snaA and snaB genes, corresponding to the polypeptides SnaA and SnaB, participates in the final step of biosynthesis of the II component of pristinamycins (conversion of pristinamycin IIB to pristinamycin IIA), corresponding to the oxidation of the 2,3 bond of D-proline. These two polypeptides constitute the two subunits of pristinamycin IIA synthase, the purification of which is described in the present invention. The product of the samS gene is considered to participate in the synthesis of SAM (methyl group donor) from ATP and methionine. The A component of most streptogramins is, in effect, methylated at C-4 (Figure 1), and this methyl has been described (Kingston *et al.*, 1983) as being derived from the methyl of methionine, very probably via a methylation reaction with SAM. The samS gene is hence considered to code for a SAM synthase (SamS; EC. 2.5.1.6) which is specific to the biosynthesis pathway of pristinamycins.

The snbA, snbR, papA and papM genes are present on cosmid pIBV2 (Figure 5). The snbA gene corresponds, on the basis of the biochemical studies presented in Example 5, to the first step for synthesis of pristinamycins I. This comprises activation of the first acid of the chain, 3-hydroxypicolinic acid, by adenylation. The snbR gene might participate in the transport of molecules of the pristinamycin I (or

possibly pristinamycin II) family out of the cell after synthesis, thereby conferring a resistance to this component on the producing strain. The papA gene corresponds, on the basis of sequence analyses (Example 5 8.8) and the study of a mutant disrupted in this gene (Example 9.3), to a gene for the biosynthesis of para-aminophenylalanine from chorismate. para-Aminophenylalanine is then dimethylated by the product of the papM gene, an N-methyltransferase described in the present 10 invention, to form para-dimethylaminophenylalanine, which is then incorporated in pristinamycin IA. The papA and papM genes hence participate in the synthesis of one of the precursors of pristinamycin IA.

The snaA, snaD, snbC, snbD and snbE genes are 15 present on cosmid pIBV3 (Figure 6), which hence adjoins cosmid pIBV1 on which the snaA gene is already present. The snaD gene codes, on the basis of analysis of its sequence (Example 8.9) and the study of a mutant disrupted in this gene (Example 9.5), for a peptide 20 synthase involved in the biosynthesis of pristinamycin II. The snbC gene, whose product is described in the present invention, participates in the incorporation of threonine and aminobutyric acid residues in the peptide chain of pristinamycin IA. The snbD gene, whose product 25 is also described in the present invention, is involved in the incorporation of proline and para-dimethylaminophenylalanine residues in the peptide chain of pristinamycin IA. It also governs the

N-methylation of the peptide bond between these 2 residues. Lastly, the snbE gene, whose product is also described in the present invention, participates in the incorporation of the last two residues of pristinamycin 5 IA, namely phenylglycine and 4-oxopipeolic acid.

The snaC gene is present on cosmid pIBV4 (Figure 7). It codes for an FMN:NADH oxidoreductase, also designated FMN reductase, described in the present invention and which supplies pristinamycin IIA synthase 10 with FMNH₂ from FMN and NADH. The snaC gene hence participates in the final step of the biosynthesis of pristinamycin IIA.

These different genes were subcloned from their cosmid of origin and their nucleic acid sequences 15 were determined. The snaA, snaB and samS genes were subcloned on a 6-kb BamHI-BamHI fragment, a portion of which was sequenced (SEQ ID no. 1). The snbA gene was subcloned in a 5.5-kb EcoRI-BglII fragment, a portion of which was sequenced (SEQ ID no. 5). The snbR gene 20 was subcloned in a 4.6-kb BglII-BglII fragment, a portion of which was sequenced (SEQ ID no. 6). A portion of the papA gene was subcloned in a 3.4-kb XhoI-XhoI fragment, a portion of which was sequenced (SEQ ID no. 9). The papM gene was subcloned in a 4.1-kb 25 PstI-PstI fragment, a portion of which was sequenced (SEQ ID no. 10). A portion of the snaD gene was subcloned in a 1.5-kb BamHI-SstI fragment, a portion of which was sequenced (SEQ ID no. 8). A portion of the

snbC gene was subcloned on a 6.2-kb SphI-SphI fragment, 2 regions of which were sequenced (SEQ ID nos. 11 and 12). A portion of the snbD gene was subcloned on an 8.4-kb SphI-SphI fragment, 2 regions of which were 5 sequenced (SEQ ID Nos. 13 and 14). A portion of the snbE gene was subcloned on a 6.6-kb SphI-SphI fragment, 2 regions of which were sequenced (SEQ ID Nos. 15 and 16). The snaC gene was subcloned in a 4-kb BamHI-BamHI fragment, a portion of which was sequenced (SEQ ID 10 no. 7).

The proximity of the snaA, snaB, snaD, samS, snbC, snbD and snbE genes on the one hand, as well as the snbA, snbR, papA and papM genes, confirms the cluster localization of the genes for biosynthesis of 15 the A and B components of streptogramins. Furthermore, the 4 cosmids described in the present invention are grouped together in a region of the chromosome whose size is estimated at 200 kb by pulsed-field 20 electrophoresis, equivalent to 3% of the total genome (7500 kb) of Streptomyces pristinaespiralis (Example 13). It is hence obvious that the regions surrounding the genes identified in the present invention (snaA, snaB, snaD, samS, snbC, snbD and snbE; snbA, snbR, papA and papM; snaC) contain the other genes of the 25 pristinamycin biosynthesis cluster, and that these genes may be used to localize the other genes for the biosynthesis of streptogramins.

Preferably, the subject of the invention is a

nucleotide sequence chosen from:

(a) all or part of the snaA (SEQ ID no. 2), snaB (SEQ ID no. 3), snaC (SEQ ID no. 7), snaD (SEQ ID no. 8), papA (SEQ ID no. 9), papM (SEQ ID no. 10), sams (SEQ ID no. 4), snbA (SEQ ID no. 5), snbC (SEQ ID nos. 11 and 12), snbD (SEQ ID nos. 13 and 14), snbE (SEQ ID nos. 15 and 16) and snbR (SEQ ID no. 6) genes,

10 (b) the sequences adjacent to the genes (a) constituting the biosynthesis clusters and coding for the polypeptides involved in the biosynthesis of streptogramins,

15 (c) the sequences which hybridize with all or part of the genes (a) or (b) and which code for a polypeptide involved in the biosynthesis of streptogramins, and

(d) the sequences derived from the sequences (a), (b) and (c) owing to the degeneracy of the genetic code.

20 Still more preferably, the subject of the invention is the nucleotide sequences represented by the snaA (SEQ ID no. 2), snaB (SEQ ID no. 3), snaC (SEQ ID no. 7), snaD (SEQ ID no. 8), papA (SEQ ID no. 9), papM (SEQ ID no. 10), sams (SEQ ID no. 4), snbA (SEQ ID no. 5), snbC (SEQ ID nos. 11 and 12), snbD (SEQ ID nos. 13 and 14), snbE (SEQ ID nos. 15 and 16) and snbR (SEQ ID no. 6) genes.

25 Another subject of the invention relates to any recombinant DNA comprising a gene for the

biosynthesis of streptogramins. More preferably, this is a recombinant DNA comprising all or part of cosmids pIBV1, pIBV2, pIBV3 or pIBV4 as shown in Figures 4 to 7, or all or part of sequences which hybridize with 5 cosmids pIBV1 to pIBV4 or with fragments of these latter.

In a preferred embodiment of the invention, the nucleotide sequences defined above form part of an expression vector, which can be autonomously 10 replicating or integrative.

As stated above, although the invention is more especially illustrated with the genes for the biosynthesis of pristinamycin, it is clear that the results obtained apply to all streptogramins.

15 More especially, the techniques developed in the present invention for purifying proteins or cloning genes for the biosynthesis of streptogramins from S. pristinaespiralis may be applied to other microorganisms producing streptogramins (see Table 1).

20 Thus, the purification of an enzymatic activity from S. pristinaespiralis makes it possible to purify the same activity from another strain producing streptogramin. The present invention may hence by applied to the cloning of genes for the biosynthesis of 25 streptogramins from any producing microorganism, by purification of a protein participating in the biosynthesis and then, using the NH₂-terminal sequence thereof, synthesis of an oligonucleotide probe which

enables the corresponding gene to be cloned. Chromosome walking then enables the whole biosynthesis cluster to be identified.

Furthermore, from the genes identified in the present application, it is possible, by hybridization, to clone the genes for the biosynthesis of streptogramins directly from the DNA of another producing microorganism. In effect, the genes for the biosynthesis of pristinamycins hybridize strongly with those for the other streptogramins. It is thus possible to clone, by hybridization, the genes for the biosynthesis of streptogramins using as a probe the sna, snb or pap genes, or fragments of the latter, or fragments adjacent to these containing, as is shown in the present invention, other sna and snb genes. This is due to the fact that: 1) the streptogramins produced by the different microorganisms have identical or similar structures (see Figure 3), 2) the genes for the biosynthesis of streptogramins are organized in clusters, and 3) the enzyme systems responsible for this biosynthesis do not have an absolute specificity for their substrates.

Moreover, the cloning of genes involved in the biosynthesis of streptogramins may also be carried out using degenerate oligonucleotides, prepared from the sequences of the sna or snb genes mentioned above, or fragments of these genes, or fragments adjacent to these genes. It is thus possible to take one's pick of

the genes for the biosynthesis of the A and B components of the different strains producing streptogramins. These strains can belong to the genus Streptomyces, and also to other genera (see Table 1).

5 In addition, if the genomic DNA of the starting strains used has a G+C composition different from that observed in Streptomyces, the probes used may be synthesized with a codon bias specific to the genus or species from which it is desired to isolate the DNA.

10 Another subject of the present invention relates to the polypeptides resulting from the expression of the nucleotide sequences defined above. More especially, the present invention relates to polypeptides comprising all or part of the polypeptides 15 SnaA (SEQ ID no. 2), SnaB (SEQ ID no. 3), SnaC (SEQ ID no. 7), SnaD (SEQ ID no. 8), PapA (SEQ ID no. 9), PapM (SEQ ID no. 10), SamS (SEQ ID no. 4), SnbA (SEQ ID no. 5), SnbC (SEQ ID nos. 11 and 12), SnbD (SEQ ID nos. 13 and 14), SnbE (SEQ ID no. 15 and 16) and SnbR (SEQ ID 20 no. 6) or of derivatives of these. Within the meaning used in the present invention, the term derivative denotes any molecule obtained by modification of a genetic and/or chemical nature of the peptide sequence. Modification of a genetic and/or chemical nature is 25 understood to mean any mutation, substitution, deletion, addition and/or modification of one or more residues. Such derivatives may be generated for different purposes, such as, in particular, that of

increasing the affinity of the peptide for its substrate(s), that of improving its levels of production, that of increasing its resistance to proteases, that of increasing and/or modifying its 5 activity, or that of endowing it with novel biological properties. Among derivatives resulting from an addition, there may be mentioned, for example, chimeric polypeptides containing an additional heterologous portion attached to one end. The term derivative also 10 comprises polypeptides homologous to the polypeptides described in the present invention and originating from other cell sources, and in particular from strains producing streptogramins.

The subject of the invention is also any 15 recombinant cell containing a nucleotide sequence or a vector as defined above. The recombinant cells according to the invention can equally well be eukaryotic cells or prokaryotic cells. Among eukaryotic cells which are suitable, animal cells, yeasts or fungi 20 may be mentioned. In particular, as regards yeasts, yeasts of the genus Saccharomyces, Kluyveromyces, Pichia, Schwanniomyces or Hansenula may be mentioned. As regards animal cells, COS, CHO, C127 cells, Xenopus eggs, and the like, may be mentioned. Among fungi, 25 special mention may be made of Micromonospora, Aspergillus ssp. or Trichoderma ssp. As prokaryotic cells, it is preferable to use the following bacteria: Actinomycetes, and Streptomyces in particular, E. coli

(Example 11), Bacillus. Preferably, the recombinant cells of the invention are chosen from cells producing streptogramins (see Table 1). The recombinant cells of the invention may be obtained by any method which 5 enables a foreign nucleotide sequence to be introduced into a cell. It can be, in particular, transformation, electroporation, conjugation, protoplast fusion or any other technique known to a person skilled in the art.

A further subject of the invention is a 10 method for producing a polypeptide involved in the biosynthesis of streptogramins, according to which a recombinant cell as defined above is cultured and the polypeptide produced is recovered.

The subject of the invention is also the use 15 of a recombinant cell as defined above, expressing at least one polypeptide involved in the biosynthesis of streptogramins, in a bioconversion reaction. In particular, these cells can enable a streptogramin to be converted into a derived form. For example, 20 pristinamycin IIB can be converted in this manner to pristinamycin IIA. The same reasoning may be applied to any biosynthesis intermediate. These cells can also enable hybrid antibiotics having advantageous pharmacological properties to be manufactured (Hopwood 25 et al. 1985a, Hopwood et al. 1985b, Hutchinson et al. 1989). These bioconversions may be carried out either using whole cells, or using acellular extracts of the said cells.

Another subject of the invention relates to the use of a nucleotide sequence as defined above for amplifying streptogramin production. The invention also relates to a method for producing streptogramins, 5 according to which one or more nucleotide sequences according to the invention is/are introduced and/or amplified in a cell producing streptogramins or which is potentially a producer of streptogramins, the said cell is cultured under conditions of streptogramin 10 production, and the streptogramins produced are recovered.

The overexpression of certain genes involved in the biosynthesis can enable the streptogramin A and/or B production of the producing strains to be 15 increased. This overproduction may be carried out in several strains: either strains which produce only molecules of the streptogramin A family, or strains which produce only molecules of the streptogramin B family, or strains which produce both the A and B 20 components. These overexpressions can result from an increase in the level of synthesis, and hence in the productivity, of the A and/or B components, either in an Erlenmeyer, or in small fermenters, or in large industrial fermenters. Moreover, the specific 25 overexpression of a gene involved in the biosynthesis of an A or B component also makes it possible to vary the % of A and B components produced by the strain, and thus to obtain a better synergy between these

molecules. In addition, the biosynthesis genes isolated from a microorganism producing streptogramins may be used to amplify production in another producing microorganism.

5 Another subject of the invention relates to a method for preparing cells blocked in a step of the pathway of biosynthesis of streptogramins, according to which a mutagenesis is performed on at least one gene of the biosynthesis pathway, on a cell producing
10 streptogramins.

Preferably, the mutagenesis is performed in vitro or in situ, by suppression, substitution, deletion and/or addition of one or more bases in the gene in question, or by gene disruption.

15 Another aspect of the present invention lies, in effect, in the construction of mutants blocked in certain steps of biosynthesis of streptogramins. The value lies, on the one hand in the study of the functionality of the mutated proteins, and on the other hand in the production of strains producing
20 biosynthesis intermediates. These intermediates may be modified, where appropriate after separation, either by adding particular components to the production media, or by introducing into the strains thus mutated other
25 genes capable of modifying the intermediate by acting as a substrate for them. These intermediates may thus be modified by chemical, biochemical, enzymatic and/or microbiological means. In this context, the mutant

SP92::pVRC505 of S. pristinaespiralis strain SP92 was constructed: S. pristinaespiralis SP92::pVRC505 was isolated by homologous integration in the snaA gene of a suicide plasmid pVRC505, constructed from the vector 5 pDH5 and a fragment internal to the snaA gene. The following mutants were also constructed: SP92 samS::QamR; SP92::pVRC508; SP92::pVRC404 and SP92::pVRC1000 (Example 9).

The invention hence also relates to a method 10 for preparing an intermediate of the biosynthesis of streptogramins, according to which:

- a cell blocked in a step of the pathway of biosynthesis of streptogramins is prepared as described above,
- 15 - the said cell is cultured, and
- the accumulated intermediate is recovered.

The invention also relates to a method for preparing a molecule derived from streptogramins, according to which:

20 - a cell blocked in a step of the pathway of biosynthesis of streptogramins is prepared as described above,

- the said cell is cultured, and
- the intermediate accumulated by this cell 25 is modified, where appropriate after separation of the culture medium.

The present invention is illustrated by means of the examples which follow, which are to be

considered as illustrative and non-limiting.

LIST OF FIGURES

Figure 1: Example of structure of the A components of streptogramins.

5 Figure 2: Example of structure of the B components of streptogramins.

Figure 3: Other examples of structures of streptogramins.

Figure 4: Diagram of cosmid pIBV1.

10 Figure 5: Diagram of cosmid pIBV2.

Figure 6: Diagram of cosmid pIBV3.

Figure 7: Diagram of cosmid pIBV4.

Figure 8: Reaction catalysed by pristinamycin IIA synthase.

15 Figure 9: Reaction catalysed by 3-hydroxypicolinic acid:AMP ligase.

Figure 10: Reaction catalysed by SnbC.

Figure 11: Reaction catalysed by SnbD.

Figure 12: Reaction catalysed by SnbE.

20 Figure 13: Reaction catalysed by SnaC.

Figure 14: Reaction catalysed by PapM.

Figure 15: Diagram of plasmids pVRC402 (A) and pVRC501 (B).

Figure 16: Diagram of plasmid pXL2045.

25 Figure 17: Diagram of plasmid pVRC1105.

Figure 18: Diagram of plasmid pVRC1106.

Figure 19: Diagram of plasmid pVRC1104.

Figure 20: Diagram of plasmid pVRC900.

Figure 21: Diagram of plasmid pVRC1000.

Figure 22: Diagram of plasmid pVRC509.

Figure 23: Diagram of plasmid pVRC903.

Figure 24: Diagram of plasmid pVRC409.

5 Figure 25: Diagram of plasmid pVRC505.

Figure 26: Diagram of plasmid pVRC701.

Figure 27: Diagram of plasmid pVRC702.

Figure 28: Diagram of plasmid pVRC508.

Figure 29: Diagram of plasmid pVRC404.

10 Figure 30: Diagram of plasmid pVRC507.

Figure 31: Diagram of plasmid pVRC706.

Figure 32: General map.

MATERIALS

Bio-Sil SEC 125 and 250 columns (Bio-Rad)

15 MonoQ HR 5/5, 10/10 and 16/10 columns
(Pharmacia)

PD-10 column (Pharmacia)

Superose 6 HR 10/30 column (Pharmacia)

Superdex 200 Hi-Load 16/60 and 75 HR 10/30

20 column (Pharmacia)

Superose 12 prep grade column (Pharmacia)

Vydac C4 and C18 columns (The Separations
Group)

Nucleosil 5-C18 column (Macherey-Nagel)

25 Phenyl Superose HR 10/10 column (Pharmacia)

TSK G2000 SW column (Tosoh, Japan)

Phenyl Sepharose (Pharmacia)

FMN-agarose (Sigma)

Q Sepharose Fast Flow (Pharmacia)

Sephadex G-25 Fine (Pharmacia)

Centricon 10 or 30 (Amicon)

5 Centriprep 10 or 30 (Amicon)

Centrilutor (Amicon)

EXAMPLE 1: Isolation of total DNA of

Streptomyces pristinaespiralis strain SP92

This example illustrates how S.

10 pristinaespiralis SP92 DNA may be purified.

S. pristinaespiralis strain SP92 is derived from S. pristinaespiralis strain DS5647 (ATCC25486).

50 ml of YEME medium (34% sucrose, 5 mM MgCl₂, 0.25% glycine (D. Hopwood *et al.* 1985)) are inoculated 15 with 10⁸ S. pristinaespiralis SP92 spores, and the culture is incubated for 40 hours at 30°C with stirring at 280 rpm.

The mycelium is harvested and washed with 15 ml of 10.3% sucrose. Approximately 1 g of the 20 mycelium pellet is taken up with 5 ml of TE supplemented with 34% of sucrose, to which are added 1 ml of lysozyme at a concentration of 50 mg/ml in 10 mM Tris-HCl solution pH 8.0 and 1 ml of 0.25 M EDTA pH 8.0. After incubation at 30°C for a period of 30 to 25 60 min, the mixture is clarified by adding 0.8 ml of 10% sarkosyl. 2 ml of 0.25 M EDTA pH 8.0, 10 ml of TE, 18 g of CsCl and 1.2 ml of ETB at a concentration 10 mg/ml are then added. The preparation is

ultracentrifuged overnight at 55,000 rpm at 20°C.

The chromosomal DNA, present in the CsCl gradient in the form of a band, is recovered using a Pasteur pipette. The ETB is removed by several washes 5 with a solution of isopropanol saturated with TE buffer, 5 M NaCl. The DNA is precipitated by adding 3 volumes of TE and 4 volumes of isopropanol. After washing with 70% ethanol, the DNA is taken up in a suitable volume of TE. The total amount of DNA obtained 10 varies between 250 and 500 µg per g of mycelium.

EXAMPLE 2: Isolation of E. coli plasmid DNA

This example illustrates how E. coli plasmid DNA is prepared from recombinant strains of E. coli.

2.1. Preparation of E.coli plasmid DNA in 15 large amounts

This example illustrates how maxi preparations of plasmid DNA are produced in E. coli.

This preparation is performed using a 500 ml culture in LB medium containing 150 µg/ml of 20 ampicillin. The extraction protocol is derived from the methods described by Birnboim and Doly (1979) and Ish-Horowicz and Burke (1981), and is described in Maniatis et al. (1989).

After this extraction, the plasmid DNA is 25 purified using a CsCl gradient as described by Maniatis et al. (1989). The plasmid DNA is then precipitated by adding 3 volumes of TE and 4 volumes of isopropanol. After centrifugation, the pellet is taken up in 0.5 to

1 ml of TE.

2.2. Preparation of *E. coli* plasmid DNA in small amounts

This example illustrates how minipreparations 5 of plasmid DNA are produced in *E. coli*.

This preparation is carried out using 1.5 ml of culture in LB medium containing 150 µg/ml of ampicillin. The procedure is that described by Birnboim and Doly (1979).

10 EXAMPLE 3: Construction of the genomic DNA library of *S. pristinaespiralis* SP92 in *E. coli* and preparation of hybridization membranes

This example illustrates how a genomic DNA library of *S. pristinaespiralis* SP92 is produced in *E. coli*. 15

3.1. Preparation of genomic DNA fragments

This example illustrates how high molecular weight genomic DNA fragments may be prepared.

Total DNA of the strain SP92, prepared as 20 described in Example 1, is partially digested with Sau3A (New England Biolabs, Beverly, MA. 01915-5510 USA) in the buffer recommended by the supplier: 100 mM NaCl, 10 mM Tris-HCl (pH7.5), 10 mM MgCl₂, 100 µg/ml BSA. The amount of enzyme used to obtain high molecular 25 weight DNA fragments was determined empirically.

Approximately 0.025 enzyme units are used to digest 1 µg of total DNA for 20 min at 37°C. The reaction is then stopped by incubation for 15 min at 65°C, and the

enzyme is removed by adding an equal volume of phenol/chloroform. After centrifugation, the supernatant containing the partially digested total DNA is precipitated by adding 0.3 M final sodium acetate and 5 2.5 volumes of ethanol.

Approximately 100 µg of total DNA are digested in this way, and DNA fragments between 30 and 50 kb in size are isolated with a 10-40% sucrose gradient. Their size is verified by electrophoresis on 10 0.4% agarose gel.

3.2. Preparation of cosmid pHC79

This example illustrates how cosmid pHC79 is prepared from E. coli.

Cosmid pHC79 (Hohn, B. and Collins, 1980) 15 comprises a portion of pBR322 (Bolivar, F. et al., 1977), the cro-cII region of λ and the region containing the cos sequence of Charon 4A (Blattner, F.R. et al., 1977).

Extraction of the cosmid was carried out as 20 described in Example 2.1., from an E. coli strain TG1 (K12, Δ (lac-pro) supE thi hsd DS F' traD36 proA^{B+} lacI^q LacZ Δ M15, Gibson, 1984).

500 ng of cosmid pHC79 are digested with BamH I (New England Biolabs, Beverly, MA. 01915-5510 25 USA) in 20 µl of buffer comprising 150 mM NaCl, 6 mM Tris-HCl pH 7.9, 6 mM MgCl₂, 6 mM 2-mercaptoethanol, 100 µg/ml BSA.

3.3. Ligation of the DNA fragments and the cosmid

This example illustrates how the fragments of the S. pristinaespiralis SP92 genome originating from 5 an Sau3A digestion may be ligated with the BamHI- linearized vector pHC79.

Approximately 150 ng of cosmid linearized as described above were precipitated by means of ethanol with 350 ng of fragments of total DNA of S.

10 pristinaespiralis SP92 prepared as described in Example 3.2. The pellet was taken up in 10 μ l of ligation buffer: 50 mM Tris-HCl pH 7.8, 10 mM MgCl₂, 20 mM DTT, 1 Mm ATP, 50 μ g/ml of BSA, and 0.5 μ l of T4 DNA ligase at a concentration of 400,000 units per ml (New England 15 Biolabs, Beverly, MA. 01915-5510 USA) were added. Incubation was carried out overnight at 15°C.

3.4. Carrying out encapsidation in vitro

This example illustrates how the cosmids constructed in 3.3 are encapsidated in vitro.

20 Encapsidation of the hybrid cosmids after ligation was carried out using the Gigapack II Gold kit developed by Stratagene (Stratagene Cloning Systems, La Jolla, CA 92037, USA).

25 2 \times 4 μ l of ligation mixture, equivalent to 2 \times 70 ng of hybrid cosmids, were encapsidated in vitro according to the procedure described by the supplier.

3.5. Transfection of *E. coli* strains DH1 and HB101

This example illustrates how the cosmids are introduced into *E. coli*.

5 Two transfections were carried out in parallel with *E. coli* strains DH1 (F^- gyrA96 recA1 relA1 endA1 thi-1 hsdR17 supE44L-, Low 1968) and HB101 (F^- supE44 hsdS20(rB⁻mB⁻) recA13 ara-14 proA2 lacY1 galK2 rpsL20 xyl-5 mtl-1, Boyer and Roulland-Dussoix 1969).

10 The cells were prepared according to the following protocol: a 100-ml preculture is produced in LB medium supplemented with 0.2% maltose and 10 mM MgSO₄, for 4 to 5 hours until the OD₆₀₀ reaches a value of 0.8. The culture is then centrifuged, and the pellet is 15 taken up in 40 ml of 10 mM MgSO₄, and diluted to OD₆₀₀ = 0.5 in the same solution. 200 μ l of the cell suspension thus prepared are mixed with 100 μ l of encapsidation mixture. After 20 min of contact at 37°C, 1 ml of LB is added and the whole is incubated for 20 1 hour at 37°C. The transfectants are then selected on solid LB medium containing 150 μ g/ml of ampicillin. The number of transfectants obtained is approximately 10⁴ per μ g of recombinant cosmid.

3.6. Storage of genomic DNA libraries of *S.*

25 *pristicaespiralis* SP92

This example illustrates how the genomic DNA libraries of *S. pristicaespiralis* SP92 are stored.

After verification of the average size of the

fragments inserted into cosmid pHC79, approximately 1500 colonies originating from each of the transfections carried out with the strains HB101 and DH1 are subcultured in 96-well microtitration plates 5 containing 200 μ l of Hogness medium (LB medium supplemented with 8.8% glycerol, 3 mM sodium acetate, 55 mM K_2HPO_4 , 26 mM KH_2PO_4 , 1 mM $MgSO_4$, 15 mM $(NH_4)_2SO_4$, 150 μ g/ml ampicillin). These plates are incubated overnight at 37°C and then stored at -80°C.

10 3.7. Preparation of hybridization membranes from genomic libraries of *S. pristinaespiralis* SP92

This example illustrates how the DNA of the colonies constituting the genomic libraries of *S. pristinaespiralis* SP92 is transferred onto a 15 hybridization membrane.

These hybridization membranes were produced in duplicate for each of the 2 libraries according to the following protocol:

The 15 microtitration plates of each library 20 are replicated using a replica plater on LB agar medium containing 150 μ g/ml of ampicillin. After growth overnight at 37°C, colony transfer is performed onto a Biohylon-Z⁺ membrane (Bioprobe System) according to the following protocol: the membrane is cut to the 25 appropriate size and left in contact with the colonies for 1 min. Denaturation is then performed by soaking the membrane with 0.5 M NaOH, 1.5 M NaCl solution for 5 min, followed by neutralization by soaking the

membrane in 3 M sodium acetate solution for 5 min. The DNA is fixed to the membrane by exposure under a UV lamp for 5 min.

EXAMPLE 4

5

4.1. Preparation of chromosomal DNA of *S. pristinaespiralis* strain SP92 and strains derived from SP92 in the form of inserts for pulsed-field electrophoresis

This example illustrates how DNA of S. pristinaespiralis strain SP92 and strains derived from SP92 is prepared in the form of inserts for pulsed-field electrophoresis.

This preparation is made from a mycelium culture obtained in the following manner: 30 ml of YEME medium containing 0.25% of glycine are inoculated with 10⁸ spores of the strain under study, and the culture is incubated for 48 hours at 30°C and stirred at 280 rpm in 250-ml Erlenmeyers. The mycelium is then harvested by centrifugation for 10 min at 3800 rpm and washed twice with 10% sucrose. The mycelium pellet is then resuspended in 5 ml of solution I (250 mM EDTA pH 8.0, 20.6% sucrose). To 200 Ml of mycelium thereby obtained, 400 Ml of a lysozyme solution at a concentration of 50 mg/ml in solution I together with 800 Ml of 1% LMP agarose in 25 mM EDTA pH 8 and 10.3 % sucrose, maintained at 42°C, are added. The mixture maintained at 42°C is then poured into the wells of special combs, which are closed with adhesive tape and kept for 30 min

at 4°C. The mixture solidifies, and the 30 to 40 inserts thereby obtained and contained in the wells are carefully removed from the moulds.

The inserts are first rinsed for 30 min at 5 4°C in a solution containing 25 mM EDTA and 10.3% sucrose. They are then soaked in a solution of 500 mM EDTA, 1% lauryl sarcosyl and 1 mg/ml of proteinase K for twice 24 hours at 50°C, stirring from time to time. The inserts are then washed for 3 times one hour in TE 10 containing 1 mM PMSF, changing the solution after each wash. The inserts thereby obtained are stored at 4°C for not more than 4 months in 0.5 M EDTA pH 8.0.

4.2. Digestion of inserts of DNA of *S. pristinaespiralis* strain SP92 and strains derived from 15 SP92 and analysis by pulsed-field electrophoresis

This example illustrates how chromosomal DNA of *S. pristinaespiralis* strain SP92 and strains derived from SP92, prepared in the form of inserts as described in Example 4.1., is cut with different restriction 20 enzymes for pulsed-field electrophoresis.

4.2.1. Digestion of chromosomal DNA in the form of inserts:

The inserts are first washed six times in TE, and then incubated twice for one hour in the buffer of 25 the chosen restriction enzyme. Each insert is then placed in the lid of an Eppendorf tube containing 160 Ml of buffer of the restriction enzyme and 40 units of enzyme. The whole is covered with Parafilm, and the

Eppendorf is closed to hold in place the Parafilm which enables any evaporation of the buffer to be avoided. The tubes are incubated at the desired temperature in an incubator overnight.

5 4.2.2. Analysis of digested DNA by pulsed-field electrophoresis:

The pulsed-field electrophoresis technique chosen for this study is that of the CHEF (Clamped Homogenous Electric Field) system developed by Chu et 10 al. (1986), which makes it possible to obtain two homogeneous alternating fields oriented at 120° with respect to one another and linear trajectories for the DNA molecules. The apparatus used is the "Pulsafor System" marketed by Pharmacia-LKB.

15 The electrophoretic migration parameters, such as the pulse time and the migration period, were varied so as to obtain an optimal separation of DNA fragments ranging in size between 10 and 2500 kb. The three migration conditions used are as follows: to 20 separate large fragments from 200 to 1700 kb in size, the chosen migration is 40 hours with a pulse time of 90 seconds; to separate fragments from 50 to 400 kb in size, the chosen migration is 20 hours with a pulse time of 10 seconds followed by 20 hours with a pulse 25 time of 30 seconds; lastly, to separate smaller fragments from 10 kb to 200 kb in size, the chosen migration is 24 hours with a pulse time of 10 seconds. For these three migration conditions, the voltage is

set at a constant 150 volts, the temperature is maintained at 13°C and the electrophoresis gels contain 1.3% of agarose.

The inserts containing chromosomal DNA of 5 S. pristinaespiralis strain SP92 and strains derived from SP92 are digested with the restriction enzymes as described above and are placed in the wells of the electrophoresis gel using two scalpel blades. The molecular weight markers used are "Yeast chromosome PFG 10 marker" and "Lambda Ladder PFG marker" marketed by the company New England Biolabs. Migration is performed under one of the conditions described above and the gel is then stained in a bath of ETB (ethidium bromide) at a concentration of 4 Mg/ml for 20 min and thereafter 15 decolorized in water for 20 min. After the gel is photographed, the DNA fragments are transferred onto a nylon membrane and then hybridized with [α -³²P]dCTP-labelled probes as described in Example 9.1.

EXAMPLE 5: Isolation of cosmids carrying the 20 genes coding for purified proteins involved in the biosynthesis of streptogramins

This example describes how, starting from a purified protein participating in biosynthesis of pristinamycins and whose NH₂-terminal sequence or an 25 internal sequence has been established, it is possible to isolate a cosmid carrying the structural gene for this same protein from the genomic libraries produced above, or alternatively to identify the corresponding

structural gene from among the genes carried by the cosmids and which have already been sequenced.

5.1. Isolation of cosmids pIBV1 and pIBV3
carrying one or both structural genes for the two

5 subunits of pristinamycin IIA synthase

5.1.1. Identification and purification of one of the proteins involved in the final step of the synthesis of pristinamycins II: pristinamycin IIA synthase

10 As stated in the introduction, the final step of synthesis of pristinamycin IIA corresponds to an oxidation of the 2,3 bond of D-proline to dehydroproline. The protein responsible for this activity has been purified to homogeneity, as
15 illustrated by this example.

5.1.1.A. Assay of pristinamycin IIA synthase activity

This example illustrates the assay of an activity of the biosynthesis pathway of pristinamycin 20 IIA which has never before been described and which possesses the noteworthy property of being expressed only during the period of production of pristinamycins. The enzyme in question is pristinamycin IIA synthase, which catalyses the conversion of pristinamycin IIB to 25 pristinamycin IIA by oxidation of the D-proline residue of pristinamycin IIB to a 2,3-dehydroproline residue (Figure 8) in the presence of molecular oxygen and FMNH_2 . The enzyme fractions to be assayed (0.002 to

0.005 units) are incubated for 1 h at 27°C in a total volume of 500 ml of 50 mM bis-tris propane buffer pH 6.8 containing NADH (500 µM), FMN (5 µM), pristinamycin IIB (20 µM) and 0.02 units of FMN 5 reductase (Boehringer Mannheim).

The pristinamycin IIA formed is assayed by HPLC after incubation is stopped by adding 500 µl of 0.1 N hydrochloric acid and 500 µl of acetonitrile and centrifugation of the sample for 5 min at 5000 g.

10 150 µl of the centrifugation supernatant are injected onto a 15-cm Nucleosil 5-C8 column eluted with a mixture of 34% of acetonitrile and 66% of 0.1 M phosphate buffer pH 2.9. Pristinamycins IIA and IIB are detected by means of their UV absorbance at 206 nm.

15 The unit of enzymatic activity is defined as the amount of enzyme needed to synthesize 1 µmol of pristinamycin IIA per hour under the conditions described.

5.1.1.B. Purification of S. pristinaespiralis

20 SP92 pristinamycin IIA synthase

This experiment illustrates how an enzyme of S. pristinaespiralis SP92 participating in the biosynthesis pathway of pristinamycin IIA may be purified.

25 Using the assay described above in Example 5.1.1.A, the purification of pristinamycin IIA synthase is carried out as described below taking care to freeze and store the active fractions at -30°C between

successive steps if necessary.

150 g of a centrifugation pellet, washed with 0.1 M phosphate buffer pH 7.2 containing 10% v/v of glycerol, of an S. pristinaespiralis SP92 culture

5 harvested at the beginning of the pristinamycin production phase are taken up with 450 ml of 50 mM bis-tris propane buffer pH 6.8 containing 5 mM DTT and 0.2 mg/ml of lysozyme. The suspension thereby obtained is incubated for 45 minutes at 27°C and then

10 centrifuged at 50,000 g for 1 hour. The crude extract thereby collected is fractionated by ammonium sulphate precipitation. The protein fraction precipitating at between 40 and 55% saturation is desalted on a column of Sephadex G-25 Fine, and then injected (100 mg per

15 injection) in pH 6.8 50 mM bis-tris propane buffer, 1 mM DTT onto a monoQ HR 10/10 column. The proteins are eluted with a linear KCl gradient (0 to 0.5 M). The fractions containing the enzymatic activity (detected by means of the test described in Example 5.1.1.A) are

20 pooled and concentrated to 20 ml on Centriprep 10.

After dilution with one volume of pH 6.8 50 mM bis-tris propane buffer, 1 mM DTT containing 2 M ammonium sulphate, the proteins are chromatographed (22.5 mg per injection) on a Phenyl Superose HR 10/10 column with a

25 decreasing ammonium sulphate gradient (1.0 M to 0 M).

The best fractions containing the desired activity are pooled, reconcentrated to 1 ml on Centriprep 10 and then applied (200 µl per injection) to a Bio-Sil

SEC 250 column. The activity peak is detected in this technique at a molecular weight centred at 77,000. The fraction containing the activity is injected onto a MonoQ HR 5/5 column in pH 6.8 50 mM bis-tris propane buffer, DTT 1 mM eluted with a linear KCl gradient (0 to 0.5 M).

After this step, the enzyme is pure and, in SDS-PAGE electrophoresis, two subunits of molecular weight estimated at 35,000 and 50,000 are detected.

They are separated on a 25-cm Vydac C4 column eluted with a linear gradient of from 30 to 50% of acetonitrile in water containing 0.07% of trifluoroacetic acid.

Table: Purification of pristinamycin IIA synthase

15	Purification step	Vol. (ml)	Protein (mg)	Sp.Act. $\mu\text{mol/h/mg}$	Yield (%)	Purification factor
	Crude extract	490	1690	0.14	100	1
	40-45% A.S.	60	1050	0.19	85	1.4
	MonoQ 10/10	95	45	3.0	58	21
20	Phenyl Superose	8	2.8	12	14	86
	Bio-Sil SEC	5	1.3	18	14	130
	MonoQ 5/5	10	0.7	23	10	160

The purification factor is calculated from the increase in specific activity of the fractions during the purification.

5.1.2. Production of oligonucleotides from

the protein sequences:

This example describes how, starting from the NH₂-terminal sequences of the two subunits of pristinamycin IIA synthase purified as described in

5 Example 5.1.1.B., it is possible to synthesize oligonucleotides. The two subunits of pristinamycin IIA synthase are referred to as SnaA and SnaB, and correspond to polypeptides of molecular weights 50,000 and 35,000, respectively, as described in Example

10 5.1.1.B.

The NH₂-terminal sequences of the proteins SnaA and SnaB, corresponding to the subunits of pristinamycin IIA synthase, were deduced by microsequencing. This is carried out by the Edman

15 degradation technique, using an automated sequencer (Applied Biosystems model 407A) coupled to an HPLC apparatus for identification of the phenylthiohydantoin derivatives. About thirty residues were determined for each of them.

20 Protein SnaA: (see residues 2 to 29 on SEQ ID No. 2)

T A P(R)(R,W)R I T L A G I I D G P G G H V A A(W)R H P (A) T

Protein SnaB: (see residues 2 to 31 on SEQ ID No. 3)

T A P I L V A T L D T R G P A A T L G T I T(R)A V(R)A A E A

Moreover, sequences internal to these two

25 polypeptides were determined after trypsin digestion of SnaA and SnaB and purification of the fragments obtained on a Vydac C18 HPLC column. The following internal sequences were found:

Protein SnaA: (see residues 365 to 384 on SEQ ID No. 2)

G A D G F N I D F P Y L P G S A D D F V

Protein SnaB: (see residues 122 to 136 on SEQ ID No. 3)

G L(-)D S F D D D A F V H D R

5 From the underlined regions in each of the sequences of the fragments internal to the proteins SnaA and SnaB, and in accordance with the degeneracy of the genetic code specific to Streptomyces (see Example 8), the following mixtures of oligonucleotides were 10 synthesized with a Biosearch 8600 automated synthesizer. They were then purified by the technique already described (Sawadogo M. and Von Dyke M. W., 1991). The snaA and snaB genes denote the structural genes for the proteins SnaA and SnaB, respectively.

15 Mixture corresponding to the underlined portion of the internal sequence of SnaA:

ATC	GAC	TTC	CCC	TAC	CTC	CCC	GG
T	T	G	T	G	G		
				A			
20				T			

Mixture corresponding to underlined portion of the internal sequence of SnaB:

TTC	GAC	GAT	GAT	GCA	TTC	GTC	CAT	GAC
C	C	T	C	G	G	C	C	

5.1.3. Labelling of the mixtures of synthetic oligonucleotides and hybridization with the genomic DNA libraries of the strain SP92

30 This example describes how oligonucleotides specific for a gene for the biosynthesis of

pristinamycins may be radioactively labelled and then hybridized with membranes onto which DNA of genomic libraries of S. pristinaespiralis SP92 has been transferred.

5 Labelling of the oligonucleotides is carried out by transfer at the 5'-terminal position of the [γ - 32 P]phosphate group of ATP with T4 polynucleotide kinase. This labelling is carried out as described in Maniatis et al. (1989). After labelling, the 10 oligonucleotides are used without purification.

Approximately 2 \times 500 ng of each mixture of oligonucleotides were labelled in this way with 32 P and were used to hybridize each of the two libraries.

15 Hybridization of the membranes of each library is carried out according to a protocol derived from those developed by Meinkoth, J. and Wahl, G. (1984) and Hames, B.D. and Higgins, S.J. (1985): the 15 membranes are prehybridized for 3 hours at 50°C in 40 ml of a solution containing: Denhardt (x5) [Denhardt 20 (x100): 2% (w/v) Ficoll, 2% (w/v) polyvinylpyrrolidone, 2% (w/v) BSA)], SSC (x5) [SSC (x20): 3 M NaCl, 0.3 M sodium citrate], 50 mM NaPO₄, pH 6.5, 0.1% SDS, 250 μ g/ml salmon sperm DNA].

25 Hybridization is then carried out overnight at 50°C in 20 ml of the same solution to which the 500 ng of labelled oligonucleotides are added.

The filters are then washed in a solution of SSC (x6) and 0.5% SDS, twice for 30 min at room

temperature and then empirically at gradually higher temperatures (50 to 65°C). The temperature of these latter washes is gradually increased after successive autoradiographic exposures in order to determine the 5 specificity of the hybridizing clones with the mixtures of oligonucleotides.

5.1.4. Isolation of cosmids pIBV1 and pIBV3 and determination of the regions containing the snaA and snaB genes

10 This example illustrates how it is possible to isolate cosmids constructed as described in Example 3 containing genes for the biosynthesis of pristinamycins.

15 Cosmids pIBV1 and pIBV3 were isolated from two clones originating, respectively, from the library produced in the strain HB101 and from the library produced in the strain DH1 which hybridized with both mixtures of oligonucleotides simultaneously for pIBV1 and with the mixture of oligonucleotides originating 20 from the internal sequence of the protein SnaA for pIBV3.

These cosmids were purified as described in Example 2. Cosmids pIBV1 and pIBV3 contain, respectively, a genomic DNA insert of S. 25 pristinaespiralis SP92 whose sizes were estimated, respectively, at 30 kb and 34 kb. Maps (Figures 4 and 6) were established from digestions with different restriction enzymes, according to the protocols of the

supplier (New England Biolabs, Beverly, MA. 01915-5510 USA).

Southern hybridizations of pIBV1 and pIBV3 DNA, digested by means of different enzymes, with the 5 mixtures of oligonucleotides enabled the region of this cosmid containing the snaA and/or snaB genes to be identified.

Southern hybridization was carried out as described in Maniatis et al. (1989). After separation 10 of the restriction fragments by electrophoresis on 0.8% agarose gel, the DNA is transferred onto a Biohylon Z⁺ membrane (Bioprobe System). Hybridization of the DNA thus transferred onto the membranes with the mixtures of oligonucleotides was carried out as described in 15 Example 5.1.3.

These Southern hybridizations enabled it to be shown that cosmid pIBV1 possessed a 6-kb BamHI fragment containing the sequences homologous to the probes synthesized in Example 5.1.2 (originating from 20 the proteins SnaA and SnaB), as well as a 2.5-kb EcoRI fragment internal to the BamHI fragment containing the sequences homologous to the probes originating exclusively from the protein SnaA. Furthermore, the hybridization signals obtained with cosmid pIBV3 showed 25 that it possessed only the 2.5-kb EcoRI fragment containing the sequences homologous to probes originating exclusively from the protein SnaA.

5.2. Isolation of cosmid pIBV2 containing the structural gene for 3-hydroxypicolinic acid:AMP ligase (snbA)

This example illustrates how it is possible
5 to obtain a cosmid as constructed in Example 3
containing at least one gene for the biosynthesis of
pristinamycins I.

5.2.1. Identification and purification of the protein involved in the activation of 3-hydroxypicolinic acid

10

This example illustrates how the protein
responsible for the activation of 3-hydroxypicolinic
acid may be purified to homogeneity from S.
pristinaespiralis SP92.

15 5.2.1.A. Assay of 3-hydroxypicolinic acid:AMP ligase

This example illustrates the assay of an
activity of the biosynthesis pathway of pristinamycin
IA which has never before been described and which
20 possesses the noteworthy property of being expressed
only during the period of production of pristinamycins.
The enzyme in question is 3-hydroxypicolinic acid:AMP
ligase, which catalyses the formation of the adenylate
of 3-hydroxypicolinic acid (Figure 9) from this free
25 acid and ATP in the presence of MgCl₂.

The enzyme fractions to be assayed (0.002 to
0.020 units) are incubated for 15 min at 27°C in a
total volume of 250 µl of pH 6.8 50 mM bis-tris propane

buffer, 1 mM DTT, 10% v/v glycerol, in the presence of 3-hydroxypicolinic acid (1 mM), ATP (2 mM), MgCl₂ (5 mM) and tetrasodium pyrophosphate labelled with the radioactive isotope 32 of the phosphorus atom (200 µM).

5 The reaction is stopped by adding 1 ml of a suspension of activated charcoal at a concentration of 10 g/l in a mixture of 75% of 0.1 M tetrasodium pyrophosphate and 25% of 14% perchloric acid. After stirring, the charcoal is collected and washed with 10 twice 1 ml of the pyrophosphate/perchloric acid mixture. The radioactive organic molecules are then eluted with three times 1 ml of a mixture of 50% of methanol and 50% of N ammonia solution into a counting vial containing 12 ml of water. The radioactivity is 15 measured by the Cerenkov effect with a scintillation counter (PACKARD Minaxi TriCarb 4000).

The unit of enzymatic activity is defined as the amount of enzyme needed to incorporate 1 µmol of pyrophosphate into ATP in the course of 1 hour under 20 the conditions described above.

5.2.1.B. Purification of S. pristinaespiralis

SP92 3-hydroxypicolinic acid:AMP ligase

This experiment illustrates how an enzyme of S. pristinaespiralis SP92 participating in the 25 biosynthesis pathway of pristinamycin IA may be purified.

Using the assay described above in Example 5.2.1.A, the purification of 3-hydroxypicolinic

acid:AMP ligase is carried out as described below, taking care to freeze the active fractions at -70°C and store them at -30°C between successive steps if necessary.

5 234 g of a centrifugation pellet, washed with 0.1 M phosphate buffer pH 7.2 containing 10% v/v of glycerol, of an S. pristinaespiralis SP92 culture harvested at the beginning of the pristinamycin production phase are taken up with 234 ml of pH 8.0

10 100 mM Tris-HCl buffer containing 4 mM DTE, 1 mM benzamidine, 1 mM PMSF, 15% v/v glycerol and 0.6 mg/ml of lysozyme. The suspension thereby obtained is incubated for 30 minutes at 27°C and then centrifuged at 50,000 g for 1 hour. The crude extract thereby

15 collected is injected in pH 8.0 100 mM Tris-HCl buffer, 4 mM DTE, 1 mM benzamidine, 1 mM PMSF, 15% v/v glycerol onto a column (80 ml) of Q Sepharose Fast Flow. The proteins are eluted with a linear KCl gradient (0 to 0.4 M). The fractions containing the enzymatic activity

20 (detected by means of the test described in Example 5.2.1.A) are pooled and diluted with one volume of pH 8.0 100 mM Tris-HCl buffer, 1 mM benzamidine, 1 mM PMSF, 15% v/v glycerol containing 2 M ammonium sulphate. The proteins are then chromatographed on a

25 column (50 ml) of Phenyl Sepharose with a decreasing ammonium sulphate gradient (1.0 M to 0 M) in pH 8.0 100 mM Tris-HCl buffer, 1 mM benzamidine, 1 mM PMSF, 15% v/v glycerol. After the addition of 4 mM DTE, the

active fractions are pooled, concentrated to 5 ml on Centriprep 10 and then applied to a column (100 ml) of Superose 12 prep grade. The fractions containing the desired activity are pooled and injected in pH 8.0

5 100 mM Tris-HCl buffer, 4 mM DTE, 1 mM benzamidine, 1 mM PMSF, 15% v/v glycerol (approximately 6 mg per injection) onto a column of MonoQ HR 5/5 eluted with a linear KCl gradient (0 to 0.4 M). The active fractions are pooled, concentrated to 1 ml on Centricon 10,

10 diluted with 3 volumes of pH 6.8 50 mM bis-tris propane buffer, 4 mM DTE, 1 mM benzamidine, 1 mM PMSF, 15% v/v glycerol, and then injected (2 mg per injection) in the latter buffer onto a column of MonoQ HR 5/5 eluted with a linear KCl gradient (0 to 0.3 M). The best fractions

15 containing the desired ligase are pooled and then applied in pH 6.8 20 mM sodium phosphate buffer, 50 mM sodium sulphate to a Bio-Sil SEC 250 column. The activity peak is detected in this technique at a molecular weight centred at 60,000.

20 The protein possessing the activity of activation of 3-hydroxypicolinic acid is hereinafter designated SnbA.

..... After this step, the enzyme is pure and, in SDS-PAGE electrophoresis, its molecular weight is
25 estimated at approximately 67,000.

Table: Purification of 3-hydroxypicolinic acid:AMP ligase

Purification step	Vol. (ml)	Protein (mg)	Sp.Act. $\mu\text{mol/h/mg}^a$	Yield (%)	Purification factor
5	Crude extract	246	2050	(0.06)	
	Q Sepharose	40	188	0.47	100
	Phenyl Sepharose	70	35	2.21	4.7
	Superose 12	16	17	2.03	4.3
	MonoQ pH 8.0	4.5	9.0	2.09	4.5
	MonoQ pH 6.8	1.0	2.0	2.9	6.2
	Bio-sil 250	2.5	0.23	12.4	26

* The activity in the crude extract cannot be measured accurately owing to exchanges between pyrophosphate and ATP which are not specific to 3-hydroxypicolinic acid.

The purification factor is calculated from the increase in specific activity of the fractions during the purification.

5.2.2. Production of oligonucleotides from the protein sequence:

This example describes how, starting from the NH₂-terminal and internal sequences of the protein 3-hydroxypicolinic:AMP ligase, it is possible to synthesize oligonucleotides.

The NH₂-terminal sequence of the protein SnbA was deduced by microsequencing as described in Example

5.1.2. About twenty residues were identified in this way.

5 A sequence of approximately 20 amino acids internal to the protein SnbA was also identified after trypsin hydrolysis and purification of the fragments obtained on a Vydac C18 HPLC column.

NH₂-terminal sequence of the protein

3-hydroxypicolinic:AMP ligase:

(See residues 1 to 21 on SEQ ID No. 5)

10 M L D G S V P W P E D V A A K Y R A A G Y

Internal sequence of the protein

3-hydroxypicolinic:AMP ligase:

(See residues 448 to 467 on SEQ ID No. 5)

V S A (-) E V E G H L G A H P D V Q Q A A

15 From the underlined regions in each of the sequences, and in accordance with the degeneracy of the genetic code specific to Streptomyces (see Example 8), the following mixtures of oligonucleotides were synthesized:

20 Mixture corresponding to the underlined portion of the NH₂-terminal sequence of the protein

3-hydroxypicolinic:AMP ligase:

5'

3'

GTC CCC TGG CCC GAG GAC GTC GCC GCC AAG TAC

25 G G G G G G

Mixture corresponding to the underlined portion of the internal sequence of the protein

3-hydroxypicolinic:AMP ligase:

5'

3'

5 GAG GTC GAG GGC CAC CTC GGC GCC CAC CCC GAC GTC CAG CAG GC
 G G G G G G

5.2.3. Labelling of the mixtures of synthetic oligonucleotides and hybridization of the genomic DNA libraries of S. pristinaespiralis SP92.

10 This example describes how oligonucleotides specific for a gene for the biosynthesis of pristinamycins may be radioactively labelled and then hybridized with membranes onto which DNA of genomic libraries of S. pristinaespiralis has been transferred.

15 Labelling the oligonucleotides is carried out by transfer at the 5'-terminal position of the [γ - 32 P]phosphate group of ATP with T4 polynucleotide kinase, as described in Example 5.1.3.

20 Approximately 2 x 500 ng of each mixture of oligonucleotides were labelled in this way with 32 P and were used to hybridize each of the two libraries.

Hybridization of the membranes of each library was carried out as described in Example 5.1.3.

25 5.2.4. Isolation of cosmid pIBV2 and determination of the region containing the structural gene for 3-hydroxypicolinic acid:AMP ligase

This example illustrates how it is possible to obtain a cosmid as constructed in Example 3

containing at least the structural gene for 3-hydroxypicolinic acid:AMP ligase.

Cosmid pIBV2 was isolated from a clone of the library produced in E. coli strain DH1 which hybridized 5 with both mixtures of oligonucleotides simultaneously.

This cosmid was purified as described in Example 2. It contains a genomic DNA insert of S. pristinaespiralis SP92 whose size was estimated at 47 kb. A map (Figure 5) was established from digestions 10 with different restriction enzymes, as described in Example 5.1.4.

Southern hybridizations of pIBV2 DNA, digested by means of different enzymes, with the mixtures of oligonucleotides enabled the region 15 containing the structural gene for 3-hydroxypicolinic acid:AMP ligase to be identified. Southern blotting and hybridizations were carried out as described in Example 5.1.4.

The hybridization results enabled it to be 20 shown that cosmid pIBV2 possessed a 5.5-kb EcoRI-BglII fragment containing the sequence homologous to the probes synthesized in Example 5.2.2.

5.3. Demonstration of the presence of a portion of the structural gene for pristinamycin I 25 synthase II (SnbC) on cosmid pIBV3

This example illustrates how it is possible to identify the presence of genes for the biosynthesis of pristinamycins I on a cosmid which has already been

isolated (Example 5.1).

5.3.1. Identification of pristinamycin I synthase II involved in the incorporation of threonine and aminobutyric acid residues into the peptide chain
5 pristinamycin IA

This example illustrates how the protein responsible for the incorporation of threonine and aminobutyric acid residues into the peptide chain of pristinamycin IA may be purified to homogeneity from S.
10 pristinaespiralis SP92.

5.3.1.A. Assay of the partial activities of pristinamycin I synthase II

This example illustrates the assay of activities of the biosynthesis pathway of pristinamycin
15 IA which have never before been described and which possess the noteworthy property of being expressed only during the period of production of pristinamycins. The activities in question are the partial activities of the peptide synthase responsible for the incorporation
20 of threonine and aminobutyric acid residues into the peptide chain of pristinamycin IA (Figure 10) in the presence of ATP and MgCl₂.

25 The threonine:AMP ligase and aminobutyric acid:AMP ligase activities are measured in an enzymatic test of ATP-pyrophosphate exchange similar to that described in 5.2.1.A for 3-hydroxypicolinic acid:AMP ligase.

The aminoacylation reactions of the enzyme

with threonine or alanine (an analogue of aminobutyric acid which is found in pristinamycin IC) enable the peptide synthase to be differentiated from other enzymes which may effect an ATP-pyrophosphate exchange, 5 and in particular aminoacyl-tRNA synthetases. The test of aminoacylation of the enzyme with tritium-labelled threonine described below is hence the one which was used in this example.

10 The enzyme fractions to be assayed (0.2 to 2 units) are incubated for 15 min at 27°C in a total volume of 250 µl of pH 6.8 50 mM bis-tris propane buffer, 1 mM DTT, 10% v/v glycerol in the presence of 1 µCi of [3-³H]-L-threonine (15 Ci/mmol), ATP (2 mM) and MgCl₂ (5 mM).

15 The reaction is stopped by adding 150 µl of 25% trichloroacetic acid solution. The precipitated proteins are collected on a microfilter and washed with 3 times 400 µl of 7% trichloroacetic acid, before being eluted with twice 400 µl of N sodium hydroxide into a 20 counting vial containing 1 ml of N HCl and 12 ml of scintillation cocktail (Beckmann Readygel). The amount 25 of radioactivity contained in this vial is measured with a scintillation counter (PACKARD Minaxi TriCarb 4000). It represents the amount of threonine bound covalently to the desired peptide synthase.

The unit of enzymatic activity is defined as the amount of enzyme needed to bind 1 picomole of threonine covalently in 15 min under the conditions

described above.

5.3.1.B. Purification of pristinamycin I synthase II

This experiment illustrates how an enzyme of 5 S. pristinaespiralis SP92 participating in the biosynthesis pathway of pristinamycin IA may be purified.

Using the acid described above in Example 5.3.1.A., purification of the peptide synthase 10 responsible for the incorporation of threonine and aminobutyric acid residues into the peptide chain of pristinamycin IA is carried out as described below, taking care to work at 4°C and to store the active fractions at -70°C.

15 150 g of a centrifugation pellet, washed with 0.1 M phosphate buffer pH 7.2 containing 10% v/v of glycerol, of an S. pristinaespiralis SP92 culture harvested at the beginning of the pristinamycin production phase are taken up with 450 ml of pH 8.0 20 100 mM Tris-HCl buffer containing 4 mM DTE, 1 mM benzamidine, 1 mM PMSF, 1 mM EDTA, 1 mM EGTA, 15% v/v glycerol. The suspension thereby obtained is ground using a French Press adjusted to a pressure of 5000 psi, and then centrifuged at 50,000 g for 1 hour. 25 The crude extract thereby collected is injected in pH 8 100 mM Tris-HCl buffer, 4 mM DTE, 2 mM benzamidine, 2 mg/l leupeptin, 1 mg/l E-64, 15% v/v glycerol onto a column (200 ml) of Q Sepharose Fast Flow. The proteins

are eluted with a linear KCl gradient (0 to 0.6 M). At outflow from the column, each fraction is treated with one-tenth of its volume of a solution of 1 mM PMSF, 5 mM EDTA, 5 mM EGTA. The fractions containing the 5 enzymatic activity (detected by means of the test described in Example 5.3.1.A) are pooled and reconcentrated by ultrafiltration on Centriprep 30 to a final volume of 28 ml. This concentrate is injected in 4-ml aliquots onto a Superdex 200 Hi-Load 16/60 10 permeation column equilibrated in pH 6.8 50 mM bis-tris propane buffer, 1 mM benzamidine, 4 mM DTE, 0.2 mM Pefabloc, 1 mM EDTA, 0.1 M KCl, 20% v/v glycerol. After assaying, the active fractions are pooled and reconcentrated to 15 ml on Centriprep 30, then desalted 15 on PD-10 in pH 8.0 100 mM Tris-HCl buffer, 4 mM DTE, 2 mM benzamidine, 2 mg/l leupeptin, 1 mg/l E-64, 20% v/v glycerol and applied in two portions to a MonoQ HR 10/10 column equilibrated and eluted with a linear gradient of from 0.4 M KCl in this same buffer. The 20 fractions containing the desired activity are pooled, reconcentrated on Centriprep 30 and then Centricon 30 to a final volume of 1 ml and injected in five portions onto a column of Superose 6 HR 10/30 in pH 6.8 50 mM bis-tris propane buffer, 1 mM benzamidine, 4 mM DTE, 25 0.2 mM Pefabloc, 1 mM EDTA, 0.1 M KCl, 20% v/v glycerol. The activity peak is detected in this technique at a molecular weight centred at 450,000. After this step the enzyme is pure and, in

SDS-PAGE electrophoresis, its molecular weight is estimated at approximately 240,000. This band also contains all radioactivity of the protein labelled by aminoacylation with tritiated threonine.

5 At this stage, the maximal activity of the enzyme using a concentration of 100 μ Ci/ml of threonine (15 ci/mmol) amounts to 3670 units/mg; the enzyme is also capable of forming adenylates with L-aminobutyric acid or L-alanine; an aminoacylation reaction of the 10 enzyme with tritiated alanine is detected, and the maximal activity in the presence of 200 μ Ci/ml of [2,3-³H]-L-alanine (15 Ci/mmol) is 2290 pmol/mg in 15 min.

Table: Purification of pristinamycin I synthase II

15	Purification step	Vol. (ml)	Protein (mg)	Sp.Act.* (units/mg)	Yield (%)	Purification factor
	Crude extract	445	4700	(1)	-	-
	Q Sepharose	308	834	7	100	1
	Superdex 200	120	105	22	40	3.1
20	MonoQ HR	15	11.5	96	19	14
	Superose-6	7.5	2.8	122	6	17

a: The activity in the crude extract cannot be measured accurately.

25 The purification factor is calculated from the increase in specific activity of the fractions

during the purification.

5.3.2. Production of oligonucleotides from the protein sequence

This example describes how, starting from 5 internal sequences of pristinamycin I synthase II, it is possible to synthesize oligonucleotides.

The internal sequences of the peptide synthase which is responsible for the incorporation of threonine and aminobutyric acid residues into the 10 peptide chain of pristinamycin IA were deduced by microsequencing as described in Example 5.1.2. after trypsin hydrolysis and purification of the fragments obtained on a Vydac C18 column.

Sequences internal to the protein

15 pristinamycin I synthase II

(See residues 49 to 61 on SEQ ID No. 12)

1 5 10

L A A F N D T A R P V P R

1 5 10 15 20

20 V P A A F V P L D A L P L T G N G V L D

From the underlined regions in these sequences, and in accordance with the degeneracy of the genetic code specific to Streptomyces (see Example 8), the following mixtures of oligonucleotides were 25 synthesized:

Mixture corresponding to the underlined portion of the sequence 1 internal to the protein pristinamycin I synthase II:

Mixture corresponding to the underlined portion of sequence 2 internal to the protein pristinamycin I synthase II:

10 5' 3
TTC GTC CCC CTC GAC GCC CTC CCC CT
G G G G G G

5.3.3. Labelling of the mixtures of synthetic oligonucleotides and Southern hybridization of cosmid

15 PIBV3 DNA

This example describes how oligonucleotides specific for a gene for the biosynthesis of pristinamycins I may be radioactively labelled and then hybridized with a membrane onto which cosmid pIBV3 DNA has been transferred.

Labelling of the oligonucleotides is carried out by transfer at the 5'-terminal position of the $[\gamma-^{32}\text{P}]$ phosphate group of ATP with T4 polynucleotide kinase, as described in 5.1.3.

25 Approximately 500 ng of the mixture of
oligonucleotides were labelled in this way with ^{32}P , and
were used for Southern hybridization of pIBV3 DNA
digested with different enzymes. These hybridizations

enabled it to be shown that a portion of the structural gene for pristinamycin I synthase II was carried by cosmid pIBV3, and enabled the region containing this gene to be identified. Southern blotting and 5 hybridization were carried out as described in Example 5.1.4.

The hybridization results enabled it to be shown that cosmid pIBV3 possessed a 6.2-kb SphI fragment containing the sequence homologous to the 10 probes synthesized in Example 5.3.2.

5.4. Demonstration of the presence of a portion of the structural gene for pristinamycin I synthase III (SnbD) on cosmid pIBV3

This example illustrates how it is possible 15 to identify the presence of genes for the biosynthesis of pristinamycins I on a cosmid which has already been isolated (Example 5.1).

5.4.1. Identification of pristinamycin I synthase III involved in the incorporation of proline 20 and p-dimethylaminophenylalanine residues into the peptide chain of pristinamycin IA

This example illustrates how the protein responsible for the incorporation of proline and p-dimethylaminophenylalanine residues into the peptide 25 chain of pristinamycin IA may be purified to homogeneity from S. pristinaespiralis SP92.

5.4.1.A. Assay of partial activities of pristinamycin I synthase III

This example illustrates the assay of activities of the biosynthesis pathway of pristinamycin IA which have never before been described and which possess the noteworthy property of being expressed only 5 during the period of production of pristinamycins. The activities in question are partial activities of the peptide synthase responsible for the incorporation of proline and para-dimethylaminophenylalanine residues into the peptide chain of pristinamycin IA (Figure 11) 10 in the presence of SAM, ATP and MgCl₂.

The proline:AMP ligase and p-dimethylamino-phenylalanine:AMP ligase activities are measured in an enzymatic test of ATP-pyrophosphate exchange similar to that described in 5.2.1.A. for 3-hydroxypicolinic 15 acid:AMP ligase.

The aminoacylation reactions of the enzyme with proline and p-dimethylaminophenylalanine make it possible to differentiate the peptide synthase from other enzymes which may perform a ATP-pyrophosphate exchange, and in particular aminoacyl-tRNA synthases. 20 The same applies to the N-methylation of the α -amino function of p-dimethylaminophenylalanine acylated on the enzyme. The latter test characteristic of N-methylation is hence the one which was used in this 25 example.

The enzyme fractions to be assayed (0.2 to 2 units) are incubated for 15 min at 27°C in a total volume of 250 μ l of pH 6.8 50 mM bis-tris propane

buffer, 1 mM DTT, 10% v/v glycerol in the presence of 1 μ Ci of [methyl-³H]-SAM (15 Ci/mmol), para-dimethylamino-L-phenylalanine (1 mM), ATP (2 mM) and MgCl₂ (5 mM).

5 The reaction is stopped by adding 150 μ l of 25% trichloroacetic acid solution. The precipitated proteins are collected on a microfilter and washed with 3 times 400 μ l of 7% trichloroacetic acid, before being eluted with twice 400 μ l of N sodium hydroxide into a
10 counting vial containing 1 ml N HCl and 12 ml of scintillation cocktail (Beckmann Readygel). The amount of radioactivity contained in this vial is measured with a scintillation counter (PACKARD Minaxi TriCarb 4000). It represents the amount of N-methylated para-
15 dimethylaminophenylalanine bound covalently to the desired peptide synthase.

The unit of enzymatic activity is defined as the amount of enzyme needed to bind 1 picomole of N-methylated p-dimethylaminophenylalanine covalently in
20 15 min under the conditions described above.

5.4.1.B. Purification of pristinamycin I synthase III

-----> This experiment illustrates how an enzyme of S. pristinaespiralis SP92 participating in the
25 biosynthesis pathway of pristinamycin IA may be purified.

Using the assay described above in Example
5.4.1.A, purification of the peptide synthase

responsible for the incorporation of proline and para-dimethylaminophenylalanine residues into the peptide chain of pristinamycin IA is carried out as described below, taking care to work at 4°C and to store the 5 active fractions at -70°C.

250 g of a centrifugation pellet, washed with 0.1 M phosphate buffer pH 7.2, 1 mM PMSF, 5 mM EDTA, 5 mM EGTA, 0.5 M KCl, 10% v/v glycerol, of an S. pristinaespiralis SP92 culture harvested at the 10 beginning of the pristinamycin production phase are taken up with 750 ml of pH 8.0 100 mM Tris-HCl buffer containing 4 mM DTE, 5 mM benzamidine, 0.2 mM Pefabloc, 1 mM EDTA, 1 mM EGTA, 2 mg/l leupeptin, 2 mg/l STI, 2 mg/l aprotinin, 1 mg/l E-64, 20% v/v glycerol. The 15 suspension thereby obtained is ground using a French Press adjusted to a pressure of 5000 psi, and then centrifuged at 50,000 g for 1 h. The crude extract thereby collected is fractionated by ammonium sulphate precipitation. The protein fraction coming out at 20 between 0 and 35% ammonium sulphate saturation is redissolved in the disruption buffer and desalted on a column of Sephadex G 25 Fine equilibrated and eluted in this same buffer. The proteins thus prepared are injected in pH 8.0 100 mM Tris-HCl buffer, 4 mM DTE, 25 2 mM benzamidine, 2 mg/l leupeptin, 1 mg/l E-64, 20% v/v glycerol onto a column (200 ml) of Q Sepharose Fast Flow, and are then eluted with a linear KCl gradient (0 to 0.6 M). At outflow from the column, each fraction is

treated with one-tenth of its volume of a solution of 2 mM Pefabloc, 5 mM EDTA, 5 mM EGTA, 5 mM benzamidine. The fractions containing the enzymatic activity (detected by means of the test described in Example 5 5.4.1.A) are pooled and precipitated with ammonium sulphate at 80% saturation. The proteins which have come out are redissolved in pH 6.8 50 mM bis-tris propane buffer, 1 mM benzamidine, 1 mM DTE, 0.2 mM Pefabloc, 1 mM EDTA, 1 mM EGTA, 2 mg/l leupeptin, 10 0.15 M NaCl, 20% v/v glycerol, and injected in 5 4-ml aliquot portions onto a Superdex 200 Hi-Load 16/60 permeation column equilibrated and eluted in this same buffer. After assay, the active fractions are pooled and reconcentrated to 3 ml on Centriprep 30, then 15 rediluted to 20 ml with pH 8.0 100 mM Tris-HCl buffer, 4 mM DTE, 1 mM benzamidine, 1 mM PMSF, 20% v/v glycerol and applied in two portions to a MonoQ HR 10/10 column equilibrated and eluted with a linear gradient from 0.4 M KCl in this same buffer. The best fractions 20 containing the desired activity are pooled and used as material for characterization of the activities of the enzyme and for its microsequencing.

After this step, the enzyme is pure and, in SDS-PAGE electrophoresis, its molecular weight is 25 estimated at approximately 250,000. This band also contains all the radioactivity of the protein labelled by aminoacylation with tritiated SAM and para-dimethylaminophenylalanine. In permeation on Superose

6 HR 10/30, the native molecular weight of the enzyme is estimated at 700,000.

At this stage, the enzyme is also capable of forming adenylates with proline; an aminoacylation reaction of the enzyme with tritiated proline is detected, and the maximal activity in the presence of 200 μ Ci/ml of [5-³H]-L-proline (34 Ci/mmol) is 2490 pmol/mg in 15 min.

Table: Purification of pristinamycin I synthase III

Purification step	Vol. (ml)	Protein (mg)	Sp.Act.* (units/mg)	Yield (%)	Purification factor
Crude extract	800	8100	(4)	-	-
35% A.S.	200	4000	(6)	-	-
Q Sepharose	132	498	46	100	1
Superdex 200	45	39.5	417	71	9
MonoQ HR	9	5.3	1070	25	23

a: The activity in the crude extract and after ammonium sulphate precipitation cannot be measured accurately.

20 The purification factor is calculated from the increase in specific activity of the fractions during the purification.

5.4.2. Production of oligonucleotides from the protein sequence

25 This example describes how, starting from

internal sequences of pristinamycin I synthase III, it is possible to synthesize oligonucleotides.

An internal sequence of the peptide synthase responsible for the incorporation of proline and para-dimethylaminophenylalanine residues into the peptide chain of pristinamycin IA was deduced by micro-sequencing as described in Example 5.1.2. after cyanogen bromide treatment and purification of the fragments obtained on a Vydac C18 HPLC column.

10 Sequence internal to the protein

pristinamycin I synthase III

1 (see residues 2 to 20 on SEQ ID No. 13)

1 5 10 15 20

P- V T P Y R A Y A L A H L A G - D D D

15 From the underlined region in this sequence, and in accordance with the degeneracy of the genetic code specific to Streptomyces (see Example 8), the following mixture of oligonucleotides was synthesized:

Mixture corresponding to the underlined

20 portion of the sequence internal to the protein

pristinamycin I synthase III:

5' 3'

— GTC ACC CCG TAC CGC GCC TAC

G G C G G

25 5.4.3. Labelling of the mixtures of synthetic oligonucleotides and Southern hybridization of cosmid pIBV3 DNA

This example describes how oligonucleotides

specific for a gene for the biosynthesis of pristinamycins I may be radioactively labelled and then hybridized with a membrane onto which cosmid pIBV3 DNA has been transferred.

5 Labelling of the oligonucleotides is carried out by transfer at the 5'-terminal position of the [γ - 32 P]phosphate group of ATP with T4 polynucleotide kinase, as described in 5.1.3.

10 Approximately 500 ng of the mixture of oligonucleotides were labelled in this way with 32 P, and were used for Southern hybridization of pIBV3 DNA digested with different enzymes. These hybridizations enabled it to be shown that a portion of the structural gene for pristinamycin I synthase III was carried by 15 cosmid pIBV3, and enabled the region containing this gene to be identified. Southern blotting and hybridization were carried out as described in Example 5.1.4.

20 The hybridization results enabled it to be shown that cosmid pIBV3 possessed an 8.4-kb SphI fragment containing the sequence homologous to the probes synthesized in Example 5.4.2.

25 5.5. Demonstration of the presence of a portion of the structural gene for pristinamycin I synthase IV (SnbE) on cosmid pIBV3

This example illustrates how it is possible to identify the presence of genes for the biosynthesis of pristinamycins I on a cosmid which has already been

isolated (Example 5.1).

5.5.1. Identification of the peptide synthase (referred to as pristinamycin I synthase IV) responsible for the incorporation of the phenylglycine residue into the peptide chain of pristinamycin IA

5.5.1.A. Assay of enzymatic activities carried by the peptide synthase (pristinamycin I synthase IV) responsible for the incorporation of the phenylglycine residue into the peptide chain of pristinamycin IA

This example illustrates the assay of an enzymatic activity of the biosynthesis pathway of pristinamycin IA which has not been described hitherto and which possesses the noteworthy property of being expressed only during the period of production of pristinamycins in the wild-type microorganism. The activity in question is that of the peptide synthase (pristinamycin I synthase IV) responsible for the incorporation of the L-phenylglycine residue into the peptide chain (Figure 12) in the presence of ATP and $MgCl_2$. The phenylglycine:AMP ligase activity of pristinamycin I synthase IV is measured in an enzymatic test of ATP-pyrophosphate exchange similar to that described in 5.2.1.A. for 3-hydroxypicolinic acid:AMP ligase activity, in the presence of L-phenylglycine (1 mM) and KCl (50 mM) in the incubation buffer.

5.5.1.B. Purification of the peptide synthase responsible for the incorporation of the phenylglycine

residue (pristinamycin I synthase IV) into the peptide chain of pristinamycin IA

This example illustrates how an enzyme of S. pristinaespiralis SP92 participating in the

5 biosynthesis pathway of pristinamycin IA may be purified. Using the assay described above in Example 5.5.1.A. The purification of pristinamycin I synthase IV is carried out as described below. All the operations are performed at 4°C. The fractions 10 containing the activity are frozen immediately and stored at -70°C.

70 g of wet cells, harvested as described in Example 5.2.1.B., are resuspended in 250 ml of cell lysis buffer (100 mM Tris-HCl pH 8.0 containing 25% of 15 glycerol, 4 mM DTE, 1 mM EGTA, 1 mM EDTA, 1 mM PMSF, 1 mg/l E-64, 2 mg/l STI, 2 mg/l α -macroglobulin, 1 mg/l leupeptin, 2 mg/l aprotinin, 5 mM benzamidine, 0.6 mg/ml lysozyme. The solution thereby obtained is kept stirring at 4°C for 1 h and then centrifuged at 20 50,000 g for 1 h. The supernatant is then injected in the cell lysis buffer onto a column of Sephadex G-25, and the excluded fraction (approximately 250 mg of protein injected in each chromatographic run) is injected onto a column of Mono Q HR 16/10 (Pharmacia) 25 equilibrated with 100 mM Tris-HCl buffer pH 8.0, 4 mM DTE, 1 mM EGTA, 1 mM EDTA, 1 mg/l E-64, 2 mg/l STI, 20% glycerol. The proteins are eluted with a linear gradient of from 0 to 0.6 M KCl and, at outflow from

the column, each fraction is treated with one-tenth of its volume of a solution of 2 mM Pefabloc, 5 mM EGTA, 5 mM EDTA. The fractions containing the activity are pooled and then mixed with 1 volume of 100 mM Tris-HCl 5 pH 8.0, 15% glycerol, 1 mM PMSF, 1 mM benzamidine, 4 mM DTT, 3.4 M ammonium sulphate per 3 volumes of fraction. The solution is injected onto a column of Phenyl Superose HR 10/10 (one-fifth of the solution is injected at each chromatographic run), and the proteins 10 are eluted with a decreasing linear gradient of from 0.9 to 0 M ammonium sulphate. The fractions containing the activity are pooled. The solution is concentrated to 3500 μ l in a Centriprep 30 and injected in two portions onto a Superdex 200 Hi-Load 16/60 column 15 equilibrated and eluted with 50 mM bis-tris propane buffer pH 6.8 containing 20% of glycerol, 0.15 M NaCl, 4 mM DTT, 1 mM PMSF, 1 mM benzamidine, 1 mM EDTA. The active fraction is diluted with 9 volumes of 50 mM bis-tris propane buffer pH 6.8 containing 25% of glycerol, 4 mM DTT, 1 mM PMSF, 1 mM benzamidine, and 20 then injected onto a column of Mono Q HR 5/5 equilibrated in the same buffer. The desired activity is eluted with a linear gradient of from 0 to 0.4 M KCl and concentrated to 630 μ l in a Centricon-30. The 25 desired protein is then purified by electrophoresis on 6% polyacrylamide gel after denaturation of the sample by heating for 10 min at 80°C with an SDS/mercapto-ethanol mixture. After electrophoresis and staining of

the gel with Coomassie blue, the gel band containing the protein is cut out and the protein is electroeluted from the gel in a Centrilutor.

Note: the band corresponding to pristinamycin

5 I synthase IV is identified by comparison with a tritiated (by covalent binding to tritiated phenylglycine; see description in Example 5.5.2.) pristinamycin I synthase IV standard.

10 After this step, the enzyme is pure in electrophoresis (SDS-PAGE). Its molecular weight is estimated at approximately 170,000.

5.5.2. Labelling of pristinamycin I synthase IV by thioesterification of radioactive phenylglycine on the enzyme

15 After activation in the form of an adenylate through phenylglycine:AMP ligase activity, phenylglycine is transferred to a thiol group of the active site of the enzyme before being incorporated into the peptide chain during elongation (general process of 20 biosynthesis of peptide antibiotics known by the name of "thiotemplate mechanism"). Generally speaking, radioactive labelling of the protein effecting the activation of amino acid may hence be performed by preparing the thioester derivative with a radioactive 25 form of the amino acid.

As an example, the radioactive labelling of pristinamycin I synthase IV is accomplished by incubating 50 µg of the protein (active fraction

emerging from the Mono Q HR 5/5 chromatography column; see above in Example 5.5.1.B.) for 1 hour at 27°C with 100 μ Ci of (RS)-2-phenyl[2- 3 H]glycine (18 Ci/mmol; Amersham) in 70 μ l of 50 mM bis-tris propane buffer pH 6.8 containing 20% of glycerol, 25 mM MgCl₂, 5 mM ATP, 0.15 M NaCl, 4 mM DTT, 1 mM PMSF, 1 mM benzamidine, 1 mM EDTA. After denaturation (SDS alone without mercaptoethanol), the proteins are separated by electrophoresis (SDS-PAGE, 6% gel) and visualized with Coomassie blue. Analysis of the radioactivity profile by counting the protein bands as well as by autoradiography (Hyperfilm MP; fluorography after impregnation of the gel with Amersham Amplify) discloses a single radioactive band with a molecular weight of 170,000.

Table: Purification of pristinamycin I synthase IV

Purification step	Protein (mg)	Sp.Act. (cpm/mg) ^a	Protein (mg)	Purification factor
Crude extract	2200	3.6	-	-
Mono Q 16/10	136	58	100	16
Phenyl Superose	32.6	175	72	49
Superdex 200	3.1	870	34	240
Mono Q 5/5	2.0	1000	25	280
Electroelution SDS-PAGE	0.1	-	-	-

*The specific activity cannot be measured accurately in the crude extract owing to the high level of non-phenylglycine-dependent ATP-pyrophosphate exchange. The specific activity value was calculated 5 from the number of units present at emergence from the first chromatographic step expressed with reference to the amount of protein in the crude extract.

5.5.3. Other activities carried by pristinamycin I synthase IV

10 Purification of the peptide synthase responsible for the incorporation of phenylglycine, described in Example 5.5.2., led to a pure protein of molecular weight 170,000. This protein does not activate the other amino acids tested, especially 15 pipecolic acid or 4-oxopipecolic acid. A second preparation of this protein, performed under the conditions described in 5.5.1.B. eliminating, however, the Phenyl Superose step, starting from another culture of S. pristinaespiralis SP92, the crude extract of 20 which was prepared in a French Press as described in 5.4.1B, led to a protein which, at emergence from the Mono Q HR 5/5 step, was equivalent in purity to that obtained at the same step in the example described in 5.5.1.B., but possessed a molecular weight of 25 approximately 250,000 in SDS-PAGE. This new preparation was competent for the activation and thioesterification of phenylglycine, but possessed, in addition, an ATP-pyrophosphate exchange activity with L-pipecolic acid

(1 mM) in the exchange test similar to that described in 5.2.1.A. for 3-hydroxypicolinic acid. Moreover, it could be shown that the 170,000 protein does not possess ATP-pyrophosphate exchange activity with L-
5 pipecolic acid even in preparations of the protein that are still very impure. It should be noted that S. pristinaespiralis SP92 naturally produces small amounts of a pristinamycin IA analogue having a pipecolic acid residue in place of 4-oxopipecolic acid. Hence this
10 demonstrates that the peptide synthase responsible for the incorporation of phenylglycine (pristinamycin I synthase IV) also catalyses the incorporation of the preceding residue (probably pipecolic acid). The difference in molecular weight obtained for
15 pristinamycin I synthase IV in the two preparations (170,000 and 250,000) is attributed to a phenomenon of partial proteolytic cleavage in the first case, leading to loss of the activity of activation of L-pipecolic acid.

20 5.5.4. Synthesis of oligonucleotides from the protein sequence

This example describes how, starting from an internal sequence of pristinamycin I synthase IV, it is possible to set about testing for the corresponding
25 gene using suitably chosen oligonucleotides.

An internal sequence of pristinamycin I synthase IV of 15 amino acids was identified after cyanogen bromide cleavage of the purified protein and

purification of the fragments obtained on a Vydac C18 HPLC column.

Sequence internal to the protein

pristinamycin I synthase IV

5 (See residues 82 to 98 on SEQ ID No. 16)

1 5 10 15

V T V F L N N T R L I O N F R P R - F - G D

From the underlined region in this sequence, and in accordance with the degeneracy of the genetic 10 code specific to Streptomyces (see Example 8), the following mixture of oligonucleotides was synthesized:

Mixture corresponding to the underlined portion of the internal sequence of the protein pristinamycin I synthase IV:

15

5' ACG CGC CTC ATC CAG AAC TTC CGC CC 3'

C	G	G	G
T			T

5.5.5. Labelling of the mixtures of synthetic 20 oligonucleotides and Southern hybridization of cosmid pIBV3 DNA

This example describes how oligonucleotides specific for a gene for the biosynthesis of pristinamycins I may be radioactively labelled and then 25 hybridized with a membrane onto which cosmid pIBV3 DNA has been transferred.

Labelling of the oligonucleotides is carried out by transfer at the 5'-terminal position of the

[γ -³²P]phosphate group of ATP with T4 polynucleotide kinase, as described in 5.1.3.

Approximately 500 ng of the mixture of oligonucleotides were labelled in this way with ³²P, and 5 were used for Southern hybridization of pIBV3 DNA digested with different enzymes. These hybridizations enabled it to be shown that a portion of the structural gene for pristinamycin I synthase II was carried by cosmid pIBV3, and enabled the region containing this 10 gene to be identified. Southern blotting and hybridization were carried out as described in Example 5.1.4.

The hybridization results enabled it to be shown that cosmid pIBV3 possessed a 6.6-kb SphI 15 fragment containing the sequence homologous to the probes synthesized in Example 5.5.4.

5.6. Isolation of cosmid pIBV4 containing the structural gene for FMN reductase (snaC)

This example illustrates how it is possible 20 to obtain a cosmid as constructed in Example 3 containing at least one gene for the biosynthesis of PII.

5.6.1. Identification of FMN reductase associated with pristinamycin IIA synthase

25 This example illustrates how the protein responsible for reduction of FMN by NADH to form the FMNH₂, needed for the reaction catalysed by pristinamycin IIA synthase may be purified to homogeneity from

S. pristinaespiralis SP92.

5.6.1.A. Assay of FMN reductase activity

This example illustrates the assay of an activity of the biosynthesis pathway of pristinamycin IIA which has never before been described and which possesses the noteworthy property of being expressed only during the period of production of pristinamycins. The enzyme in question is FMN reductase, also referred to as NADH:FMN oxidoreductase, which catalyses the reduction of FMN to FMNH₂, (Figure 13) in the presence of NADH. FMN reductases catalysing the same reaction which are specific or otherwise for NADH or NADPH and associated with other biosynthesis pathways have been described elsewhere (Duane *et al.*, 1975, Jablonski *et al.*, 1977, Watanabe *et al.*, 1982).

Two assays are used to detect this activity:

The first is based on a coupling with the pristinamycin IIA synthase described in Example 5.1.1., and is used for the first steps of the purification.

20 The enzyme fractions to be assayed (0.002 to 0.005 units) are incubated for 1 hour at 27°C in a total volume of 500 µl of 50 mM bis-tris propane buffer pH 6.8 containing NADH (500 µM), FMN (2 µM), pristinamycin IIB (20 µM) and 0.01 units of pristinamycin IIA synthase described in Example 5.1.1. The pristinamycin IIA formed is assayed by HPLC as described in Example 5.1.1.A.

The unit of enzymatic activity is defined as

the amount of enzyme needed to synthesize 1 μ mol of pristinamycin IIA per minute under the conditions described above.

The second assay is a spectrophotometric assay, and can be employed only with at least partially purified fractions. The enzyme fractions to be assayed (0.006 to 0.030 units) are incubated for 13 min at 27°C in a total volume of 3 ml of 50 mM bis-tris propane buffer pH 6.8 containing NADH (500 μ M) and FMN (2 μ M).

After 7 min of incubation, 6 readings of the optical density at 340 nm taken at 1-min intervals are performed against a reference curve without enzyme. The activity in μ mol/min is calculated by dividing the slope of decrease per min in the optical density by a factor of 6.2 (optical density of 1 mol of NADH at 340 nm).

The unit of enzymatic activity is defined as the amount of enzyme needed to consume 1 μ mol of NADH per minute under the conditions described above.

5.6.1.B. Purification of S. pristinaespiralis SP92 FMN reductase

This experiment illustrates how an enzyme of S. pristinaespiralis SP92 participating in the biosynthesis pathway of pristinamycin IIA may be purified.

Using the assays described above in Example 5.6.1.A., the purification of FMN reductase is carried out as described below, taking care to freeze and store

the active fractions at -30°C between successive steps if necessary.

500 g of a centrifugation pellet, washed with 0.1 M phosphate buffer pH 7.2 containing 10% v/v of 5 glycerol, of an S. pristinaespiralis SP92 culture harvested at the beginning of the pristinamycin production phase are taken up with 1500 ml of 50 mM bis-tris propane buffer pH 6.8 containing 5 mM DTT, 10% v/v of glycerol and 0.2 mg/ml of lysozyme. The 10 suspension thereby obtained is incubated for 45 min at 27°C and then centrifuged at 50,000 g for 1 hour. The crude extract thereby collected is fractionated by ammonium sulphate precipitation. The protein fraction precipitating at between 40 and 75% saturation is 15 desalted on a column of Sephadex G-25 Fine and then injected in pH 6.8 50 mM bis-tris propane buffer, 5 mM DTT, 10% v/v glycerol onto a column (300 ml) of Q Sepharose Fast Flow. The active proteins are not retained on the column, and they are desalted on a 20 column of Sephadex G-25 Fine and then reinjected in pH 8.2 50 mM Tris-HCl buffer, 5 mM DTT, 10% v/v glycerol onto a column (35 ml) of Q Sepharose Fast Flow and eluted with a linear KCl gradient (0 to 0.5 M). The fractions containing the enzymatic activity (detected 25 by means of the first test described in Example 5.6.1.A) are pooled, desalted on a column of Sephadex G-25 Fine and then injected in pH 8.2 50 mM Tris-HCl buffer, 5 mM DTT, 10% v/v glycerol onto a MonoQ HR

10/10 column. The proteins retained are eluted directly by the same buffer to which 0.2 M KCl has been added. They are collected in a volume of 1 ml, which is immediately reinjected onto a column of Superdex 75 HR

5 10/30 eluted with pH 6.8 50 mM bis-tris propane buffer, 1 mM DTT, 10% v/v glycerol. The fractions containing the desired activity (detected from this step onwards by means of the spectrophotometric test as described in Example 5.6.1.A) are pooled and the total volume of the

10 pool is made to 7 ml; these 7 ml are injected onto a column packed with 8 ml of FMN-agarose; the column is washed with pH 6.8 50 mM bis-tris propane buffer, 1 mM DTT, 10% v/v glycerol, and then eluted with the same buffer containing 10 μ M FMN. The active fractions are

15 pooled, desalted on PD-10 columns, injected in pH 8.2 50 mM Tris-HCl buffer, 5 mM DTT, 10% v/v glycerol onto a MonoQ HR 5/5 column and eluted with a linear KCl gradient (0 to 0.25 M).

After this step, the enzyme is pure. In SDS-
20 PAGE electrophoresis, a single fairly broad band is seen, centred at a molecular weight estimated at 28,000, while, in Bio-Sil SEC 125 gel permeation chromatography, this protein forms a symmetrical peak centred around a molecular weight of approximately 25 30,000.

For sequencing, the protein is desalted on a 25-cm Vydac C4 column eluted with a linear gradient of from 30 to 70% of acetonitrile in water containing

0.07% of trifluoroacetic acid.

Table: Purification of FMN reductase

Purification Steps	Vol. (ml)	Protein (mg)	Sp.Act. ^{a,b} (units/mg)	Yield (%)	Purification factor
5	crude extract	1620	5100	0.004 ^a	100
	40-75% A.S.	155	2930	0.005 ^a	68
	Q Seph. pH 6.8	357	180	0.058 ^a	49
	Q Seph. pH 8.2	153	15	0.36 ^a	25
10	MonoQ HR 10/10	1.0	8.8	0.50 ^a 4.4 ^b	120
	Superdex 75	1.5	3.1	7.4 ^b	200
	FMN-agarose	7.5	0.28	96 ^b	14
	MonoQ HR 5/5	3.0	0.29	68 ^b	11
	Bio-Sil 125	7.5	0.18	106 ^b	2900

a: assay coupled to pristinamycin IIA

15 synthase

b: spectrophotometric assay

The purification factor is calculated from the increase in specific activity of the fractions during the purification.

20 5.6.2. Production of oligonucleotides from the protein sequence

This example describes how, starting from NH₂-terminal and internal sequences of the protein FMN reductase, it is possible to synthesize oligo-

25 nucleotides

The NH₂-terminal sequence of FMN reductase was deduced by microsequencing as described in Example 5.1.2. About 30 residues were identified in this way.

(NH₂-Terminal sequences beginning at the 4th and at the 11th residue were also found in the sample sequenced.)

Two sequences internal to FMN reductase, of 13 and 21 amino acids, were also identified after trypsin hydrolysis and purification of fragments obtained on a Vydac C18 column.

NH₂-Terminal sequence of the protein FMN reductase

(See residues 2 to 25 on SEQ ID No. 7)

1 5 10 15 20 25
15 T G A D D P A R P A V G P Q S F R D A H A O L A S P V

Internal sequences of the protein FMN reductase:

(See residues 102 to 122 on SEQ ID No. 7)

1 5 10 15 20
20 F A G G E F A A W D G T G V P Y L P D A K

(See residues 149 to 161 on SEQ ID No. 7)

1 5 10
T G D P A K P P L L W Y R

From the underlined regions in each of the sequences, and in accordance with the degeneracy of the genetic code specific to Streptomyces (see Example 8), the following mixtures of oligonucleotides were synthesized:

Mixture corresponding to the NH₂-terminal sequence of the protein FMN reductase:

5' TTC CGC GAC GCC ATG GCC CAG CTC GC 3'
5 G G G G

Mixtures corresponding to the internal
sequences of the protein FMN reductase:

5' 3'
TTC GCC GGC GGC GAG TTC GCC GCC TGG GAC GGC ACC GG
10 G G G G G G G

5' GAC CCC GCC AAG CCC CCC CTG CTG TGG TAC CG
 G G G G C C

5.6.3. Labelling of the mixtures of synthetic oligonucleotides and hybridization of the genomic DNA libraries of *S. pristinaespiralis* SP92.

This example describes how oligonucleotides specific for a gene for the biosynthesis of pristinamycins may be radioactively labelled and then hybridized with membranes onto which DNA of genomic libraries of *S. pristinaespiralis* has been transferred

— Labelling the oligonucleotides is carried out by transfer at the 5'-terminal position of the $[\gamma-^{32}\text{P}]$ phosphate group of ATP with T4 polynucleotide kinase, as described in Example 5.1.3.

Approximately 2×500 ng of each mixture of oligonucleotides were labelled in this way with ^{32}P and

were used to hybridize each of the two libraries.

Hybridization of the membranes of each library was carried out as described in Example 5.1.3.

5.6.4. Isolation of cosmid pIBV4 and
5 determination of the region containing the structural
gene for FMN reductase (snaC)

Cosmid pIBV4 was isolated from a clone of the
library produced in E. coli strain HB101 which
hybridized with all three mixtures of oligonucleotides
10 simultaneously.

This cosmid was purified as described in
Example 2. It contains a genomic DNA insert of S.
pristinaespiralis SP92 whose size was estimated at
48 kb. A map (Figure 7) was established from digestions
15 with different restriction enzymes, as described in
5.1.4.

Southern hybridizations of pIBV4 DNA,
digested by means of different enzymes, with the
mixtures of oligonucleotides enabled the region
20 containing snaC, the structural gene for FMN reductase,
to be identified. Southern blotting and hybridizations
were carried out as described in Example 5.1.4.

25 The hybridization results enabled it to be
shown that cosmid pIBV4 possessed a 4-kb BamHI-BamHI
fragment containing the sequences homologous to the
probes synthesized in Example 5.6.3.

5.7 Demonstration of the presence of the structural gene for p-aminophenylalanine (phenyl-N)-methyltransferase on cosmid pIBV2

This example illustrates how it is possible, 5 starting from a purified protein, to identify the corresponding structural gene from among the genes which have already been analysed and sequenced as described in Examples 6.7 and 7.8 and which have also been expressed in E. coli as described in Example 11.

10 5.7.1. Identification and purification of the protein involved in the methylation of p-aminophenylalanine to p-dimethylaminophenylalanine

This example illustrates how the protein responsible for the methylation of p-aminophenylalanine 15 to p-dimethylaminophenylalanine [p-aminophenylalanine (phenyl-N)-methyltransferase] may be purified to homogeneity from S. pristinaespiralis strain SP92, and how it may also be obtained pure from a recombinant strain of E. coli.

20 5.7.1.A. Assay of the activity of methylation of p-aminophenylalanine to p-methylaminophenylalanine and of the activity of methylation of p-methylamino-phenylalanine to p-dimethylaminophenylalanine

This example illustrates the assay of two 25 terminal activities of the biosynthesis of p-dimethylaminophenylalanine, a component of pristinamycin IA. These activities have never before been described, and possess the noteworthy property of being expressed only

during the period of production of pristinamycins. They are the methylation of p-aminophenylalanine to p-methylaminophenylalanine (methylation 1) on the one hand, and the methylation of p-methylaminophenylalanine 5 to p-dimethylaminophenylalanine (methylation 2), both of these activities utilizing SAM as a methyl group donor (Figure 14).

The enzyme fractions to be assayed (1 to 20 units) are incubated for 30 min at 27°C in a total 10 volume of 200 µl of pH 6.8 50 mM bis-tris propane buffer containing SAM (200 µM) in which the methyl group is radioactively labelled with isotope 14 of the carbon atom (2 Ci/mol), in the presence of p-amino-L-phenylalanine (1 mM) for the assay of methylation 1 or 15 of p-methylamino-L-phenylalanine (2.5 mM) for the assay of methylation 2.

The reaction is stopped by adding 16 µl of 37% hydrochloric acid and then 20 µl of sodium heptane sulphonate as a concentration of 240 g/l. After 20 centrifugation, 150 µl of supernatant are injected into the HPLC system in the following gradient mode:

- mobile phase: eluent A = 1.2 g of sodium heptanesulphonate + 2.5 ml of glacial acetic acid + water
25 (qs 1000 ml)

eluent B = 1.2 g of sodium heptanesulphonate + 2.5 ml of

glacial acetic acid + 300 ml
 of acetonitrile + water
 (qs 1000 ml)

	gradient: t(min)	%B
5	0	30
	16	30
	17	100
	20	100
	21	30
10	25	30

- stationary phase: 150 x 4.6 mm Nucleosil 5 μ m
 C18 column (Macherey-Nagel)

At outflow from the column, the substrates
 and products of the enzymatic reaction are quantified
 15 by absorption at 254 nm. This detection is coupled to
 an in-line radiochemical detection by means of a
 Berthold LB506 detector equipped with a type GT400-U4
 solid scintillation cell. This enables the
 incorporation of radioactive methyl groups into the
 20 reaction products to be monitored specifically.

The unit of enzymatic activity for
 methylation 1 (for methylation 2) is defined as the
 amount of enzyme needed to incorporate 1 nmol of methyl
 groups into p-aminophenylalanine (into p-methylamino-
 25 phenylalanine).

5.7.1.B. Purification from

S. pristinaespiralis SP92 of the SAM-dependent N-methyltransferase catalysing the methylation of p-aminophenylalanine to p-dimethylaminophenylalanine

5 [p-aminophenylalanine (phenyl-N)-methyltransferase]

This experiment illustrates how an enzyme of S. pristinaespiralis SP92 participating in the biosynthesis pathway of pristinamycin IA may be purified.

10 Using the assay described above in Example 5.7.1.A, the purification of the SAM-dependent N-methyltransferase is carried out as described below, taking care to freeze and store the active fractions at -70°C between successive steps if necessary.

15 240 g of a centrifugation pellet, washed with pH 7.2 100 mM phosphate buffer, 1 mM PMSF, 5 mM EDTA, 5 mM EGTA, 0.5 M KCl, 10% v/v glycerol, of an S. pristinaespiralis SP92 culture harvested at the beginning of the pristinamycin production phase are 20 taken up in 480 ml of pH 8.0 0.1 M Tris-HCl buffer containing 4 mM DTE, 5 mM benzamidine, 0.2 mM Pefabloc, 100 µg/l E-64, 2 mg/l leupeptin, 1 mM EDTA, 1 mM EGTA, 2 mg/l STF, 2 mg/l aprotinin, 20% v/v glycerol and 2 mg/ml lysozyme, this buffer being maintained at 25 + 4°C. The suspension thereby obtained is stirred vigorously at 4°C. After 30 min of stirring, 0.2 mg/ml deoxyribonuclease I and 5 mM MgCl₂, are added. After 90 min of stirring, the extract is centrifuged for

1 hour at 50,000 g. The supernatant is divided into 3 fractions of approximately 180 ml. Each one is desalted by gel permeation on a 500 ml column of Sephadex G-25 Fine equilibrated at the natural flow rate in pH 6.8

5 20 mM bis-tris buffer containing 4 mM DTE, 5 mM benzamidine, 0.2 mM Pefabloc, 100 µg/l E-64, 2 mg/l leupeptin, 1 mM EDTA, 1 mM EGTA, 2 mg/l STI, 2 mg/l aprotinin, 20% v/v glycerol. The protein eluate is then chromatographed (400 mg of protein at each cycle) on a

10 MonoQ HR 16/10 column at a flow rate of 6 ml/min with an increasing linear gradient of sodium chloride (0 to 0.3 M) in pH 6.8 20 mM bis-tris buffer containing 4 mM DTE, 2 mM benzamidine, 100 µg/l E-64, 2 mg/l leupeptin, 20% v/v glycerol. At outflow from the column, the

15 fractions are supplemented with 10% v/v of pH 6.8 20 mM bis-tris buffer containing 4 mM DTE, 30 mM benzamidine, 2 mM Pefabloc, 100 µg/l E-64, 2 mg/l leupeptin, 5 mM EDTA, 5 mM EGTA, 10 mg/l STI, 10 mg/l aprotinin, 20% v/v glycerol. Under these conditions, both methylation

20 activities (1 and 2) are detected identically in the exclusion fractions and the first elution fractions. These fractions are pooled and concentrated by ultrafiltration on CentriPrep 10. This concentrate is made to 0.85 M ammonium sulphate and then

25 chromatographed (20 to 80 mg at each cycle) on a Phenyl Superose HR 10/10 column at a flow rate of 1 ml/min with a decreasing linear gradient of ammonium sulphate (0.85 to 0 M) in pH 6.8 50 mM bis-tris buffer

containing 4 mM DTE, 2 mM benzamidine, 100 µg/l E-64, 2 mg/l leupeptin, 1 mM EDTA, 1 mM EGTA, 10% v/v glycerol. At outflow from the column, the fractions are supplemented with 10% v/v of pH 6.8 50 mM bis-tris 5 buffer containing 4 mM DTE, 30 mM benzamidine, 2 mM Pefabloc, 100 µg/l E-64, 2 mg/l leupeptin, 1 mM EDTA, 1 mM EGTA, 10 mg/l STI, 10 mg/l aprotinin, 10% v/v glycerol. Under these conditions, both methylation activities (1 and 2) are detected identically in the 10 elution fractions corresponding to approximately 0.15 M ammonium sulphate. These fractions are pooled, concentrated by ultrafiltration on Centricon 10, desalted on PD-10 columns equilibrated in pH 8.2 (at 5°C) 50 mM Tris buffer containing 4 mM DTE, 2 mM 15 benzamidine, 100 µg/l E-64, 2 mg/l leupeptin, 20% v/v glycerol, and then chromatographed (approximately 10 mg at each cycle) on a MonoQ HR 5/5 column equilibrated in the same buffer at a flow rate of 1 ml/min. Under these conditions, the two activities are not retained on the 20 column. At outflow from the column, the exclusion fractions hence containing these two activities are supplemented with 10% v/v of pH 8.2 50 mM Tris buffer containing 4 mM DTE, 30 mM benzamidine, 2 mM Pefabloc, 100 µg/l E-64, 2 mg/l leupeptin, 1 mM EDTA, 1 mM EGTA, 25 20% v/v glycerol. These fractions are then concentrated by ultrafiltration on Centricon 10 and thereafter chromatographed on a 300 x 7.5 mm 10 µm TSK G2000 SW column equilibrated at a flow rate of 0.5 ml/min in

pH 7.0 50 mM Hepes buffer containing 4 mM DTE, 0.2 mM Pefabloc, 1 mM EDTA, 1 mM EGTA, 10% v/v glycerol, 0.15 M sodium chloride. The two activities co-elute in this technique at a retention time corresponding to a 5 molecular weight close to 30,000. After this step, a preponderant protein is visible in SDS-PAGE. It is located at around 32,000.

Table: Purification of the enzyme methylating p-aminophenylalanine to p-dimethylaminophenylalanine

10	Purification Steps	Vol. (ml)	Protein (mg)	Sp.Act. (units*/mg)	Yield (%)	Purification factor
	Crude extract	510	1800	29	-	-
	G-25 Fine	560	1560	34	102	1.17
	MonoQ HR 16/10	670	82	430	67	14.8
15	Phenyl Superose	10	3.48	6300	42	217
	MonoQ HR 5/5	7	0.88	17200	29	593
	TSK G2000	0.8	0.113	40300	8.7	1390

20 *This refers to units of enzymatic activity for methylation 1. At each step, the value of the units of enzymatic activity for methylation 2 was equal to 120% of that of the units for methylation 1.

The purification factor is calculated from the increase in specific activity of the fractions during the purification.

25 5.7.1.C. Purification from E. coli pVRC706 of the recombinant protein of S. pristinaespiralis SP92 displaying the SAM-dependent N-methyltransferase

activity catalysing the methylation of p-aminophenylalanine to p-dimethylaminophenylalanine

This experiment illustrates how an enzyme of S. prisinaespiralis SP92 participating in the 5 biosynthesis pathway of pristinamycin IA and expressed in E. coli by cloning of the papM gene may be purified.

Using the assay described above in Example 5.7.1.A., we showed that crude extracts of the recombinant strain E. coli::pVRC706 display a strong 10 activity for methylation 1 and for methylation 2, whereas in the control E. coli strain (pMTL23) neither of these two activities was detected. The purification of the SAM-dependent p-aminophenylalanine (phenyl-N)-methyltransferase catalysing the methylation of 15 p-aminophenylalanine to p-dimethylaminophenylalanine was then carried out.

Under the same conditions as those described in Example 5.7.1.B., except for a chromatography step which was eliminated (step of purification on MonoQ HR 20 5/5), we purified to homogeneity a protein which possesses a molecular weight in chromatography on a TSK G2000 column and in SDS-PAGE identical to those possessed by the protein purified in Example 5.7.1.B.

Table: Purification of the enzyme methylating
 p-aminophenylalanine to p-dimethylaminophenylalanine
 from E. coli strain pVRC706

Purification Steps	Vol. (ml)	Protein (mg)	Sp.Act. (units ^a /mg)	Yield (%)	Purification factor
crude extract	15	190	235	-	-
G-25 Fine	22	175	231	91	1
MonoQ HR 16/10	24	13.4	2100	63	8.9
Phenyl superose	3.0	0.39	35500	31	145
10 TSK G2000	0.8	0.092	45200	9.3	192

15 "This refers to units of enzymatic activity for methylation 1. At each step, the value of the units of enzymatic activity for methylation 2 was equal to 120% of that of the units for methylation 1.

15 The purification factor is calculated from the increase in specific activity of the fractions during the purification.

5.7.2. Identification of the structural gene for p-aminophenylalanine (phenyl-N)-methyltransferase

20 The NH₂-terminal sequence of the 32,000 protein purified in Example 5.7.1.B was determined by microsequencing as described in Example 5.1.2. Ten residues were determined in this way:

TAAAPTLAQAA

25 The NH₂-terminal sequence of the 32,000 protein purified in Example 5.7.1.C was determined by

microsequencing as described in Example 5.1.2. Ten residues were determined in this way:

TAAAPTLAQAA

In both cases the same residues are found,
5 and this sequence corresponds exactly to the beginning of the protein sequence which is deduced from the sequence of the papM gene (see residues 2 to 11 on SEQ ID no. 10). The purified p-aminophenylalanine (phenyl-N)-methyltransferase is hence the protein PapM.

10 EXAMPLE 6: Subcloning of DNA fragments cloned into cosmids as prepared in Example 3 and containing the genes of interest

This example illustrates how, starting from cosmids constructed as described in Example 3 and 15 containing genes for the biosynthesis of pristinamycins II or pristinamycins I, it is possible to subclone DNA fragments containing these genes.

These subclonings were performed in order to be able to deduce subsequently the nucleic acid 20 sequence of the genes identified, as well as to carry out the different construction presented in the examples which follow.

6.1. Isolation of the 5.5-kb EcoRI-BglII fragment containing the structural gene for
25 3-hydroxypicolinic acid:AMP ligase

This example describes how, starting from cosmid pIBV2 containing the structural gene for 3-hydroxypicolinic acid:AMP ligase, it is possible to

subclone a DNA fragment of smaller size containing this gene.

Approximately 10 μ g of cosmid pIBV2 were cut successively with the restriction enzymes BglII and 5 EcoRI (New England Biolabs) under the conditions recommended by the supplier. The restriction fragments thereby obtained were separated by electrophoresis on 0.8% agarose gel, and the 5.5-kb BglII-EcoRI fragment was isolated by electroelution as described in Maniatis 10 et al. (1989).

Approximately 100 ng of pUC19 (Viera and Messing 1982) cut with BamHI and EcoRI were ligated with 200 ng of the 5.5-kb BglII-EcoRI fragment under the conditions described in Example 3.3.

15 After transformation of the strain TG1 and selection of the transformants on solid LB medium containing 150 μ g/ml of ampicillin and 20 μ g/ml of X-gal according to the technique described by Maniatis 20 et al. (1989), a clone carrying the desired fragment was isolated. The recombinant plasmid was designated pVRC402. Its restriction map is presented in Figure 15(A). It was shown by hybridization, in Example 5.2., that the 5.5-kb EcoRI-BglII fragment contains the structural gene for S. pristinaespiralis SP92 25 3-hydroxypicolinic acid:AMP ligase. Plasmid pVRC402 hence contains the structural gene for S. pristinaespiralis SP92 3-hydroxypicolinic acid:AMP ligase.

6.2. Isolation of a 4.6-kb BglIII-BglIII fragment from cosmid pIBV2

This example describes how, starting from cosmid pIBV2, it is possible to subclone a DNA fragment of smaller size for the purpose of identifying, in the regions adjacent to the structural gene for 3-hydroxypicolinic acid:AMP ligase, the presence of other genes involved in the biosynthesis of pristinamycins I.

10 The different cloning steps were carried out as described above.

Approximately 10 µg of cosmid pIBV2 were cut with BglIII. The restriction fragments thereby obtained were separated by electrophoresis on 0.8% agarose gel, 15 and the 4.6-kb BglIII-BglIII fragment was isolated by electroelution.

Approximately 100 ng of pUC19 cut with BamHI were ligated with 200 ng of the BglIII-BglIII fragment.

A clone carrying the desired fragment was 20 isolated after transformation of the strain TG1 as described in Example 6.1. The recombinant plasmid was designated pVRC501. Its restriction map is presented in Figure 15(B).

6.3. Isolation of the 6-kb BamHI-BamHI fragment containing the structural genes for the two subunits of pristinamycin IIA synthase

This example describes how, starting from cosmid pIBV1, it is possible to subclone a DNA fragment

of smaller size containing the structural genes for the two subunits of pristinamycin IIA synthase.

The different cloning steps were carried out as described above.

5 Approximately 10 µg of cosmid pIBV1 were cut with BamHI. The restriction fragments thereby obtained were separated by electrophoresis on 0.8% agarose gel, and the 6-kb BamHI fragment was isolated by electroelution.

10 Approximately 100 ng of pBKS- (Stratagene Cloning Systems, La Jolla California) cut with BamHI were ligated with 200 ng of the 6-kb BamHI fragment.

A clone carrying the desired fragment was isolated after transformation of the strain TG1. The 15 recombinant plasmid was designated pXL2045. Its restriction map is presented in Figure 16. It was shown by hybridization, in Example 5.1, that the 6-kb BamHI fragment contains the snaA and snaB genes coding for the two subunits of S. pristinaespiralis SP92 20 pristinamycin IIA synthase. Plasmid pXL2045 hence contains the snaA and snaB genes coding for the two subunits of S. pristinaespiralis SP92 pristinamycin IIA synthase.

6.4. Isolation of the 6.2-kb SphI fragment

25 containing a portion of the structural gene for pristinamycin I synthase II

This example describes how, starting from cosmid pIBV3, it is possible to subclone a DNA fragment

of smaller size containing a portion of the structural gene for pristinamycin I synthase II.

The different cloning steps were carried out as described above.

5 Approximately 10 µg of cosmid pIBV3 were cut with SphI. The restriction fragments thereby obtained were separated on 0.8% agarose gel, and the 6.2-kb SphI fragment was isolated by the technique of the "Geneclean" kit marketed by the company Bio101-Ozyme.

10 Approximately 100 ng of pUC19 cut with SphI were ligated with 200 ng of the 6.2-kb SphI fragment.

A clone carrying the desired fragment was isolated after transformation of the strain TG1. The recombinant plasmid was designated pVRC1105. Its 15 restriction map is presented in Figure 17.

6.5. Isolation of the 8.4-kb SphI fragment containing a portion of the structural gene for pristinamycin I synthase III

20 This example describes how, starting from cosmid pIBV3, it is possible to subclone a DNA fragment of smaller size containing a portion of the structural gene for pristinamycin I synthase III.

The different cloning steps were carried out as described above.

25 Approximately 10 µg of cosmid pIBV3 were cut with SphI. The restriction fragments thereby obtained were separated on 0.8% agarose gel, and the 8.4-kb SphI fragment was isolated by the technique of the

"Geneclean" kit marketed by the company Bio101-Ozyme.

Approximately 100 ng of pUC19 cut with SphI were ligated with 200 ng of the 8.4-kb SphI fragment.

A clone carrying the desired fragment was 5 isolated after transformation of the strain TG1. The recombinant plasmid was designated pVRC1106. Its restriction map is represented in Figure 18.

6.6. Isolation of a 6.6-kb SphI fragment
containing a portion of the structural gene for
10 pristinamycin I synthase IV

This example describes how, starting from cosmid pIBV3, it is possible to subclone a DNA fragment of smaller size containing a portion of the structural gene for pristinamycin I synthase IV.

15 The different cloning steps were carried out as described above.

Approximately 10 µg of cosmid pIBV3 were cut with SphI. The restriction fragments thereby obtained were separated on 0.8% agarose gel, and the 6.6-kb SphI 20 fragment was isolated by the technique of the "Geneclean" kit marketed by the company Bio101-Ozyme.

Approximately 100 ng of pUC19 cut with SphI were ligated with 200 ng of the 6.6-kb SphI fragment.

A clone carrying the desired fragment was 25 isolated after transformation of the strain TG1. The recombinant plasmid was designated pVRC1104. Its restriction map is presented in Figure 19.

6.7 Isolation of the 17-kb HindIII-HindIII
fragment containing cosmid pHC79 and carrying the genes
located upstream of 3-hydroxypicolinic acid:AMP ligase
(pristinamycin I synthase I)

5 This example describes how, starting from cosmid pIBV2 containing the structural gene for 3-hydroxypicolinic acid:AMP ligase, it is possible to delete a large portion of this cosmid and retain only the portion located upstream of 3-hydroxypicolinic
10 acid:AMP ligase.

Approximately 200 ng of cosmid pIBV2 were cut with the restriction enzyme HindIII. The enzyme was denatured for 30 min at 85°C as recommended by the supplier. Cosmid pIBV2 digested in this way was
15 precipitated with ethanol as described in Maniatis et al. (1989) and religated with itself in a volume of 50 μ l.

After transformation of the strain TG1 and selection of the transformants on solid LB + 150 μ g/ml
20 of ampicillin according to the technique described by Maniatis et al. (1989), a clone containing cosmid pHC79 and the portion located upstream of 3-hydroxypicolinic acid:AMP ligase (the whole corresponding to a size of approximately 17 kb) was isolated. The recombinant
25 plasmid was designated pVRC900. Its restriction map is presented in Figure 20.

6.8. Isolation of the 1.4-kb BamHI-SstI
fragment originating from cosmid pIBV3

This example describes how, starting from cosmid pIBV3 containing the snaA gene coding for the 5 large subunit of PIIA synthase, it is possible to subclone a DNA fragment located upstream in order to study and sequence it.

Approximately 10 µg of cosmid pIBV3 were cut successively with the restriction enzymes SstI and 10 BamHI. The restriction fragments thereby obtained were separated on 0.8% agarose gel, and the 1.4-kb BamHI-SstI fragment was isolated by the technique of the "Geneclean" kit marketed by the company Bio101-Ozyme.

Approximately 100 ng of pDH5 (Hilleman et al. 15 1991) cut with BamHI and SstI were ligated with 200 ng of the BamHI-SstI fragment under the conditions described in Example 3.3.

After transformation of the strain TG1 and selection of the transformants of solid LB + 150 µg/ml 20 of ampicillin + X-gal according to the technique described by Maniatis et al. (1989), a clone carrying the desired fragment was isolated. The recombinant plasmid was designated pVRC1000. Its restriction map is represented in Figure 21.

25 6.9. Isolation of the 4-kb BamHI-BamHI
fragment containing the structural gene for FMN
reductase

This example describes how, starting from

cosmid pIBV4 containing the structural gene for FMN reductase (snaC), it is possible to subclone a DNA fragment of smaller size containing this gene.

The different cloning steps were carried out
5 as described above.

Approximately 10 µg of cosmid pIBV4 were cut with the restriction enzyme BamHI. The restriction fragments thereby obtained were separated on 0.8% agarose gel, and the 4-kb BamHI-BamHI fragment was
10 isolated by electroelution.

Approximately 100 ng of pUC19 cut with BamHI were ligated with 200 ng of the 4-kb BamHI-BamHI fragment.

After transformation of *E. coli* strain DH5 α
15 (supE44 DlacU169 (f80lacZDM15) hsdR17 recA1 endA1
gyrA96 thi-1 relA1) (Hanahan, 1983) and selection of the transformants on solid LB + 150 µg/ml of ampicillin + X-gal according to the technique described by
Maniatis et al. (1989), a clone carrying the desired
20 fragment was isolated. The recombinant plasmid was designated pVRC509. Its restriction map is presented in Figure 22.

EXAMPLE 7: Sequence of the isolated DNA fragments containing the genes for the biosynthesis of
25 pristinamycins of S. pristinaespiralis SP92

This example illustrates the sequencing of DNA fragments carrying, on the one hand genes involved in the biosynthesis of pristinamycins of the

pristinamycin I family, and on the other hand genes involved in the biosynthesis of pristinamycins of the pristinamycin II family, of the S. pristinaespiralis strain.

5 7.1. Sequencing of a 5-kb BamHI-XhoI fragment

This example illustrates how the nucleotide sequence of a fragment containing the snaA and snaB genes of S. pristinaespiralis SP92 may be obtained.

The BamHI-XhoI fragment is part of the 6-kb 10 BamHI-BamHI fragment which was cloned into phasmid pBKS- to give plasmid pXL2045 described in Example 6.3. Subfragments of this 5-kb BamHI-XhoI insert were then obtained by enzymatic digestion and thereafter 15 subcloned into phages M13mp18 or M13mp19 (Messing et al, 1981) in both orientations. The subcloning sites used are the following: EcoRI, PstI, PstI, NruI, EcoRI, NruI, NotI, SalI, SstI, XhoI, SalI and XhoI, and are shown in Figure 16.

These different inserts were sequenced by the 20 chain-termination reaction method, using as a synthetic primer the universal primer (Maniatis et al, 1989) or oligonucleotides which are synthesized (as is described in Example 5) and are complementary to a sequence of 20 nucleotides of the insert to be sequenced.

25 The overlap between these different inserts enabled the total nucleotide sequence to be established on both strands of the BamHI-XhoI fragment which comprises 5392 bp (SEQ ID no. 1).

7.2. Sequencing of a region of 1870 bp of the 5.5-kb EcoRI-BglII fragment

This example illustrates how the nucleotide sequence of a fragment containing the snbA gene of S. pristinaespiralis SP92 may be obtained.

The region of 1870 bp sequenced is part of the 5.5-kb EcoRI-BglII fragment which was cloned into plasmid pUC19 to give plasmid pVRC402 described in Example 6 (Figure 15(A)). Subfragments of the 5.5-kb EcoRI-BglII insert were obtained by enzymatic cleavage and then subcloned into phages M13mp18 or M13mp19 in both orientations. The subcloning sites are HindIII, PstI and HindIII, and are shown in Figure 15(A).

The overlap between these fragments enabled the total sequence of the Sau3A-Sau3A region, which comprises 1870 bp (SEQ ID no. 5), to be established.

7.3. Sequence of a region of 1830 bp in the 4.6-kb BglII-BglII fragment

This example illustrates how the nucleotide sequence of a fragment adjacent to that which contains the snbA gene of S. pristinaespiralis SP92 may be obtained.

.....> This sequence was deduced by subcloning the 1-kb BamHI-PstI and 2.1-kb PstI-EcoRI fragments (Figure 15(B)) from pVRC501 (Example 6) into the vectors M13mp18 and M13mp19. The PstI site was traversed by subcloning a 423-bp Sau3A-Sau3A fragment overlapping this site, followed by sequencing. The sequence of

1830 bp thereby obtained is shown in (SEQ ID no. 6).

7.4. Sequencing of two regions of 227 bp and 247 bp of the 6.2-kb SphI fragment

This example illustrates how the nucleotide sequence of fragments containing a portion of the structural gene for pristinamycin I synthase II (snbC) of S. pristinaespiralis may be obtained.

The regions of 227 and 247 bp sequenced are parts of the 6.2-kb SphI fragment which was cloned into plasmid pUC19 to give plasmid pVRC1105 described in Example 6.4 (Figure 17). Subfragments of the 6.2-kb SphI insert were obtained by enzymatic cleavage and then subcloned into phages M13mp18 or M13mp19 in both orientations. The subcloning sites are XhoI, PstI and BglII, and are shown in Figure 17. The 227-bp PstI-BglII fragment was sequenced completely, and 247 bp were sequenced from the 900-bp XhoI fragment: these sequences are presented in SEQ ID nos. 11 and 12.

7.5. Sequencing of two regions of 192 bp and 474 bp of the 8.4-kb SphI fragment

This example illustrates how the nucleotide sequence of fragments containing portions of the structural gene for pristinamycin I synthase III (snbD) of S. pristinaespiralis may be obtained.

The regions of 192 and 474 bp sequenced are parts of the 8.4-kb SphI fragment which was cloned into plasmid pUC19 to give plasmid pVRC1106 described in Example 6.5 (Figure 18). Subfragments of the 8.4-kb

SphI insert were obtained by enzymatic cleavage and then subcloned into phages M13mp18 or M13mp19 in both orientations. The subcloning sites are XhoI, PstI, SphI and BglII, and are shown in Figure 18.

5 The 192-bp BglII-SphI and 474-bp PstI-XhoI fragments were sequenced completely: these sequences are presented in SEQ ID nos. 13 and 14.

7.6 Sequencing of two regions of 485 bp and 291 bp of the 6.6-kb SphI fragment

10 This example illustrates how the nucleotide sequence of fragments containing portions of the structural gene for pristinamycin I synthase IV (snbE) of S. pristinaespiralis may be obtained.

15 The regions of 291 and 485 bp sequenced are parts of the 6.6-kb SphI fragment which was cloned into plasmid pUC19 to give plasmid pVRC1104 described in Example 6.6 (Figure 19). Subfragments of the 6.6-kb SphI insert were obtained by enzymatic cleavage and then subcloned into phages M13mp18 or M13mp19 in both 20 orientations. The subcloning sites are XhoI, PstI and SphI, and are shown in Figure 19. The 485-bp XhoI-SphI fragment was sequenced completely, and 291 bp were sequenced from the 1500-bp PstI fragment: these sequences are presented in SEQ ID nos. 15 and 16.

25 7.7. Sequence of a region of 645 bp in a 3.4-kb XhoI-XhoI fragment isolated from pVRC900

This example illustrates how the nucleotide sequence of a fragment located upstream of that which

contains the snbA gene of S. pristinaespiralis may be obtained.

To deduce this sequence, the 3.4-kb XhoI-XhoI fragment was subcloned beforehand into the vector pUC18 5 from the vector pVRC900 described in 6.7. The different cloning steps were carried out as described in 6.1: plasmid pVRC900 was digested with the restriction enzyme XhoI, and the fragments thereby obtained were separated on 0.8% agarose gel. The 3.4-kb XhoI-XhoI 10 fragment was purified by electroelution and was ligated with pUC18 cut with the restriction enzyme SalI. After transformation into TG1, a clone carrying the 3.4-kb XhoI-XhoI fragment was isolated. The recombinant plasmid was referred to as pVRC903. Its restriction map 15 is presented in Figure 23.

The 645-bp sequence was then deduced by subcloning the 1.4-kb PvuII-EcoRI and 0.9-kb PvuII-EcoRI fragments (Figure 23) from pVRC903 described above into the vectors M13mp18 and M13mp19. To carry 20 out these clonings, the vectors M13mp18 and M13mp19 were first digested with the restriction enzyme BamHI; the cohesive ends thereby liberated were filled in with the large fragment of DNA polymerase I (Klenow: New England Biolabs) according to the technique described 25 by Maniatis et al. (1989) so as to generate blunt ends compatible with the ends liberated by PvuII digestion; the vectors were then digested with the restriction enzyme EcoRI. The PvuII site was traversed by

subcloning a 2.2-kb PstI-PstI fragment, isolated from pVRC903, overlapping this site. The sequence of 645 bp thereby obtained is shown in SEQ ID no. 9.

7.8. Sequence of a region of 1050 bp in a

5 4.1-kb PstI-PstI fragment isolated from pVRC900

This example illustrates how the nucleotide sequence of a fragment located upstream of that which contains the snbA gene of S. pristinaespiralis may be obtained.

10 To deduce this sequence, a 4.1-kb PstI-PstI fragment was subcloned beforehand into the vector pUC19 from the vector pVRC900 described in 6.7. The different cloning steps were carried out as described in 6.1.

15 Plasmid pVRC900 was digested with restriction enzyme PstI, and the fragments thereby obtained were separated on 0.8% agarose gel. The 4.1-kb PstI-PstI fragment was purified by electroelution and was ligated with pUC19 cut with the restriction enzyme PstI. After transformation into TG1, a clone carrying the 4.1-kb 20 PstI-PstI fragment was isolated. The recombinant plasmid was referred to as pVRC409. Its restriction map is presented in Figure 24.

25 This sequence was then deduced by subcloning the 0.7-kb XhoI-XhoI and 1-kb XhoI-StuI fragments (Figure 24) from pVRC409 described above into the vectors M13mp18 and M13mp19. The XhoI site internal to the sequence was traversed by double-strand sequencing from plasmid pVRC409. The sequence of 1050 bp thereby

obtained is shown in SEQ ID no. 10.

7.9. Sequence of a region of 640 bp in the
1.4-kb BamHI-SstI fragment

This example illustrates how the nucleotide
5 sequence of a fragment adjacent to that which contains
the snaA and snaB genes of S. pristinaespiralis coding
for the two subunits of PIIA synthase may be obtained.

This sequence was deduced by subcloning the
1.4-kb BamHI-SstI fragment (Figure 21) from pVRC1000
10 (Example 6.8) into the vectors M13mp18 and M13mp19 (see
Example 7.1). The sequence of 640 bp obtained is shown
in SEQ ID no. 8.

7.10. Sequencing of the XhoI-KpnI region of
694 bp present in the 4-kb BamHI-BamHI fragment

15 This example illustrates how the nucleotide
sequence of a fragment containing the snaC gene of S.
pristinaespiralis may be obtained.

The region of 694 bp sequenced is part of the
4-kb BamHI-BamHI fragment which was cloned into plasmid
20 pUC19 to give plasmid pVRC509 described in Example 6.9.
A 694-bp XhoI-KpnI fragment, obtained by double
digestion of plasmid pVRC509 with the restriction
enzymes XhoI and KpnI and which hybridizes with the
3 oligonucleotide probes described in 5.6, was cloned
25 into phages M13mp18 and M13mp19. The XhoI and KpnI
subcloning sites are shown in Figure 22.

The sequence of the 694-bp fragment thereby
obtained is presented in SEQ ID no. 7.

EXAMPLE 8: Analysis of the nucleotide sequences by determination of the open reading frames

This example illustrates how it is possible to determine the open reading frames present in the 5 nucleotide sequences defined in Example 7, and to identify the genes involved in the biosynthesis of pristinamycins I and pristinamycin II of S. pristinaespiralis SP92 as well as the polypeptides encoded by these genes.

10 8.1. 5-kb BamHI-XhoI fragment (pXL2045)

This example illustrates how it is possible to determine the open reading frames present within the 5-kb BamHI-XhoI fragment isolated above and sequenced as described in Examples 6 and 7.

15 We looked for the presence of open reading frames within the 5-kb BamHI-XhoI fragment utilizing the fact that Streptomyces DNA has a high percentage of G and C bases as well as a strong bias in the use of the codons of which the coding frames are composed 20 (Bibb et al. 1984). The Staden and McLachlan (1982) method enables the probability of the coding frames to be calculated on the basis of the use of the codons of Streptomyces genes which are already sequenced and collated in a file containing 19673 codons obtained 25 from the BISANCE data-processing server (Dessen et al. 1900)..

This method enabled four highly probable open reading frames, which are shown in the following table,

to be characterized within the 5-kb BamHI-XbaI fragment. They are designated frames 1 to 4 according to their position starting from the BamHI site. For each one, their length in bases, their position within 5 the fragment (the BamHI site being located at position 1) and also the molecular weight in kDa of the corresponding protein are given. Frames 1, 3 and 4 are coded by the same strand and frame 2 by the complementary strand (Figure 16).

10	Frame number and gene name	Position	number of nucleotides	number of amino acids	MW in kDa of the protein encoded
	1 (<u>snaA</u>)	48-1313	1266	422	46.5
	2	2530- 1328	1203	401	-
15	3 (<u>snaB</u>)	(inv) 2692-	831	277	29
	4 (<u>samS</u>)	3522 3558- 4763	1206	402	43

- Frames 1 and 3 correspond respectively to the proteins SnaA (SEQ ID no. 2) and SnaB (SEQ ID no. 3) isolated above as described in Example 5 and for

which the cloning of the genes is detailed in Example 6. In effect, the NH₂-terminal sequences of the products of ORFs 1 and 3 are identical to the NH₂-terminal sequences found for the proteins SnaA and SnaB, respectively, in Example 5.1.2, apart from the amino-terminal methionine which has been excized. Moreover, the molecular masses calculated from the sequences are comparable to the apparent molecular masses of the proteins SnaA and SnaB, estimated, respectively, in SDS-PAGE as described in Example 5.

- Comparison of the product of open reading frame no. 4 with the protein sequences contained in the NBRF bank reveals a homology with various S-adenosylmethionine (or SAM) synthases, in particular of E. coli (Markham *et al.*, 1984), of rat (Horikawa *et al.*, 1989) and of S. cerevisiae (Thomas *et al.*, 1988). The percentage homology values calculated over the whole of the sequence using Kanehisa's (1984) algorithm vary from 51.8 to 55.4%.

These sequence comparisons hence enable it to be demonstrated that the product of open reading frame no. 4 is an SAM synthase involved in the biosynthesis of pristinamycins I or II. This gene was designated SamS (SEQ ID no. 4).

The demonstration of the involvement of the SamS gene in the biosynthesis of pristinamycins is confirmed by the construction of the SP92 mutant disrupted in this gene, as described in Example 9.2.

- Comparison of the sequence of the product of open reading frame no. 2 with the protein sequences contained in the Genpro bank reveals that an internal portion of this protein is 36% homologous with an 5 internal portion of the first open reading frame of the insertion sequence (IS891) of Anabaena (Bancroft and Wolk, 1989). This result suggests that open reading frame no. 2, designated ORF 401, belongs to an insertion sequence, and that there is hence an 10 insertion sequence located between the snaA and snaB genes.

8.2. 1870-bp Sau3A-Sau3A fragment (pVRC4021)

This example illustrates how it is possible to determine open reading frames present within the 15 1870-bp Sau3A-Sau3A fragment isolated above and sequenced as described in Examples 6 and 7.

The search for open reading frames for the Sau3A-Sau3A fragment was performed as above. A single complete open reading frame could be demonstrated in 20 this way. Its characteristics are as follows: this frame extends from position 109 to position 1858 of the Sau3A-Sau3A fragment, which corresponds to a frame of 1749 bases coding for a protein of 582 amino acids having a molecular mass of 61400 Da. This protein 25 corresponds to the protein SnbA purified above as described in Example 5 and for which the cloning of the gene is detailed in Example 6. In effect, the NH₂-terminal sequence of the product of the ORF present

on the Sau3A-Sau3A fragment is identical to the NH₂-terminal sequence found for the protein SnbA in Example 5.2. The molecular mass of 61400 Da calculated from the sequence is comparable to the apparent 5 molecular mass of the protein SnbA, estimated at 67000 Da in SDS-PAGE and at 60000 Da by gel permeation as described in Example 5.2.1.B.

The snbA gene hence codes for the enzyme which catalyses the formation of the acyladenylate 10 3-hydroxypicolinyl-AMP from one molecule of 3-hydroxypicolinic acid and one molecule of ATP: 3-hydroxypicolinic acid:AMP ligase (SEQ ID no. 5).

8.3. 1830-bp fragment (pVRC501)

This example illustrates how it is possible 15 to determine the open reading frames present within the 1830-bp fragment sequenced from the 3.1-kb BamHI-EcoRI fragment isolated above.

The search for open reading frames for the 1830-bp fragment was performed as above. A single 20 complete open reading frame could be demonstrated in this way. Its characteristics are as follows: the probable beginning of this frame is located at position 103 and the end at position 1686 of the region of 1830 bp sequenced from the BamHI-EcoRI fragment, which corresponds to a protein of 528 amino acids having an approximate molecular weight of 54000.

Comparison of the sequence of this protein with the sequences contained in the Genepro bank

reveals that it is homologous to proteins having a transport function for various metabolites, in particular for tetracycline in various microorganisms (Khan and Novick, 1983; Hoshino *et al.*, 1985),

5 actinorhodine (Fernandez-Moreno *et al.*, 1991) and methylenomycin (Neal and Chater, 1987b) in *S. coelicolor*.

These data indicate that the product of the open reading frame contained in the 3.1-kb BamHI-EcoRI 10 fragment is a transport protein enabling pristinamycins I (and possibly pristinamycins II) to be exported out of the cell. This protein was designated SnbR and the corresponding gene snbR (SEQ ID no. 6).

15 Analysis of the hydrophobicity profile of the protein SnbR by the method of Kyte and Doolittle (1982) corroborates its membrane localization and hence its transport function.

8.4. 1050-bp fragment (pVRC409)

This example illustrates how it is possible 20 to determine the open reading frames present within the 1050-bp fragment sequenced above from pVRC409 as described in Example 7.8.

The search for open reading frames for the 1050-bp fragment was performed as above. A single 25 complete open reading frame could be demonstrated in this way. Its characteristics are as follows: this phase extends from position 84 to position 962 of the sequenced portion, which corresponds to a frame of

878 bases coding for a protein of 292 amino acids having a molecular mass of 32000 Da. This protein was referred to as protein PapM. It was, moreover, purified from S. pristinaespiralis strain SP92 as described in Example 5. The molecular mass of 32000 Da calculated from the sequence is identical to the apparent molecular mass of 32000 Da estimated on SDS-PAGE as described in Example 5. Moreover, the NH₂-terminal sequence of this protein, deduced as described in Example 5, corresponds well to the NH₂-terminal sequence of the protein PapM identified by analysis of the open reading frames of the sequence of 1050 bp (SEQ ID no. 10).

8.5. 220-bp and 247-bp fragments (pVRC1105)

This example illustrates how it is possible to determine the open reading frames present within the 227-bp and 247-bp fragments sequenced from pVRC1105 as described in Examples 6 and 7.

The search for open reading frames for these two fragments was performed as above. An incomplete reading frame could be demonstrated in both cases over the whole length of the fragment.

The sequence obtained from the open reading frame identified on the 247-bp fragment isolated from the 900-bp XbaI fragment contains one of the internal sequences of the protein SnbC purified as described in Example 5.

Comparison of the product of the open reading

frames identified on the 227-bp and 247-bp fragments isolated from pVRC1105 with sequences of the Genpro bank reveals that they are homologous to peptide synthases. The one deduced from the 227-bp fragment 5 displays 24.5% homology with Acremonium chrysogenum (α -amino adipyl)cysteinylvaline synthetase (Gutierrez et al. 1991). The one deduced from the 247-bp fragment displays 34.9% homology with Bacillus gramicidin S synthase II (Hori et al. 1991) and 28% homology with 10 Acremonium chrysogenum (α -amino adipyl)cysteinylvaline synthetase (Gutierrez et al. 1991).

This confirms that cosmid pIBV3 isolated in Example 5.1 does indeed contain a portion of the structural gene for pristinamycin I synthase II 15 described in Example 5.3, designated SnbC (SEQ ID no. 11 and 12).

8.6. 192-bp and 474-bp fragments (pVRC1106)

This example illustrates how it is possible to determine the open reading frames present within the 20 192-bp and 474-bp fragments sequenced from pVRC1106 as described in Examples 6 and 7.

To search for open reading frames for these two fragments was performed as above. An incomplete reading frame could also be demonstrated on the 192-bp 25 fragment isolated from pVRC1106. Its characteristics are as follows: this frame begins at position 29 of the portion sequenced in the direction of BglII. No stop codon was identified, indicating that this open frame

is not terminated.

The sequence obtained from the open reading frame identified on the 192-bp BglII-SphI fragment contains the internal sequence of the protein SnbD 5 purified as described in Example 5, which proves, in fact, to be the NH₂-terminal sequence of the protein.

An incomplete reading frame could be demonstrated over the whole length of the 474-bp XbaI-PstI fragment.

10 Comparison of the product of the open reading frame identified on the 474-bp fragment isolated from pVRC1106 with the sequences of the Genpro bank reveals that this protein fragment displays from 30 to 40% homology with peptide synthases, for example 39.4% with 15 *Bacillus gramicidin S synthase II* (Hori *et al.* 1991) and 34% with *Acremonium chrysogenum* (α -amino adipyl)cysteinylvaline synthetase (Gutierrez *et al.* 1991).

20 This confirms that cosmid pIBV3 isolated in Example 5.1 does indeed contain a portion of the structural gene for pristinamycin I synthase III described in Example 5.4, designated SnbD (SEQ ID nos. 13 and 14).

8.7. 291-bp and 485-bp fragments (pVRC1104)

25 This example illustrates how it is possible to determine the open reading frames present within the 291-bp and 485-bp fragments sequenced from pVRC1104 as described in Examples 6 and 7.

The search for open frames for these two fragments was performed as above. An incomplete reading frame could be demonstrated in both cases over the whole length of the fragment.

5 The sequence obtained from the open frame identified on the 291 fragment isolated from the 1450-bp PstI fragment contains the internal sequence of the protein SnbE purified as described in Example 5.

10 Comparison of the product of the open frame identified on the 485-bp XhoI-SphI fragment isolated from pVRC1104 with the sequences of the Genpro bank reveals that it is homologous to peptide synthases, for example 34.7% homologous with *Bacillus gramicidin S synthase II* (Hori et al. 1991) and 36.2% with 15 Acremonium chrysogenum (α -aminoadipyl)cysteinylvaline synthetase (Gutierrez et al. 1991).

20 This confirms that cosmid pIBV3 isolated in Example 5.1 does indeed contain a portion of the structural gene for pristinamycin I synthase IV described in Example 5.5, designated SnbE (SEQ ID 15 and 16).

8.8. 645-bp fragment (pVRC903)

25 This example illustrates how it is possible to determine the open reading frames present within the 645-bp fragment sequenced above from plasmid pVRC903 as described in Example 6.7.

The search for open reading frames for the 645-bp fragment was performed as above. An incomplete

open reading frame could be demonstrated in this way.

Its characteristics are as follows: this frame affords two possibilities for initiation of translation, a GTG at position 61 and a GTG at position 70 of the

5 sequenced portion (the ATG located at position 124 was not taken into account owing to the sequence homologies described later). Analysis of the probabilities of the presence of Shine-Dalgarno regions does not make it possible to distinguish which of these codons

10 corresponds to the initiation. No stop codon was identified, which indicates that this open reading frame is not terminated. The gene identified in this way was referred to as papA, and the corresponding protein was referred to as protein PapA (SEQ ID no. 9).

15 Comparison of the product of the open reading frame identified in the 3.4-kb XhoI-XhoI fragment isolated from pVRC900 with sequences contained in the Genpro bank reveals that it is homologous to the II components of proteins of the p-aminobenzoate synthase 20 and anthranilate synthase type, involved, respectively, in the synthesis of p-aminobenzoic acid (folic acid precursor) and in the synthesis of anthranilic acid (tryptophan precursor) of various microorganisms. It displays, in particular, a 48% homology with the 25 protein TrpG of Azospirillum (Zimmer W., Aparicio C., and Elmerich c. Mol. Gen. Genet. (1991) 229:41-51) and a 47% homology with the protein PabA of Klebsiella pneumoniae (Kaplan J. B., Merkel W. K. and

Nichols B. P. J. Mol. Biol. (1985) 183:327-340). The proteins TrpG and PabA carry the glutaminase activity involved in the transamination of chorismic acid. The homologies demonstrated tend to show that the protein

5 PapA might be involved as well in the activity of transamination of chorismic acid. Chorismic acid is proposed as a precursor of p-dimethylaminophenylalanine, a component of pristinamycins I, by analogy with the synthesis of chloramphenicol, an antibiotic

10 deduced by Streptomyces (Sharka B., Westlake D. W. S. and Vining L. C. (1970) Chem. Zvesti 24, 66-72).

The role of the protein PapA will be shown subsequently (Example 9.3.) by analysis of mutants of the strain SP92 in the papA gene.

15 8.9. 1.5-kb BamHI-SstI fragment (PVRC1000)

This example illustrates how it is possible to determine the open reading frames present within the 1.5-kb BamHI-SstI fragment isolated above and sequenced as described in Examples 6.8 and 7.9.

20 The search for open reading frames for the sequenced region of 640 bp present in the 1.5-kb BamHI-SstI fragment was performed as described in Example 8.1. A single complete open reading frame could be demonstrated in this way. No initiation and no

25 termination of translation could be demonstrated, which indicates that the sequenced region of 640 bp is probably internal to a much larger reading frame, designated snaD (SEQ ID no. 8).

Comparison of the protein sequence encoded by the region of 640 bp with the protein sequences contained in the Genpro and NBRF banks reveals that this protein is 20-25% homologous to an internal 5 portion of peptide synthases such as B. brevis gramicidin synthase I (Hori et al. 1989), B. brevis tyrocidin synthase I (Weckermann et al. 1988) and Acremonium chrysogenum ACV synthase (Gutierrez et al. (1991)).

10 These data indicate that the protein partially encoded by the region of 640 bp is probably a peptide synthase involved in the biosynthesis of the peptide portion of pristinamycins II: in effect, all 15 the peptide synthases involved in the biosynthesis of pristinamycins I have already been identified in other regions of the S. pristinaespiralis chromosome, as is described in Examples 5.2, 5.3, 5.4 and 5.5.

8.10. 694-bp fragment (pVRC509)

20 This example illustrates how it is possible to determine the open reading frames present within the 694-bp fragment sequenced above from pVRC509 as described in Examples 6 and 7.

25 The search for open reading frames for the 694-bp fragment was performed as above. An incomplete open reading frame could be demonstrated in this way. Its characteristics are as follows: this frame begins at position 210 of the sequenced portion. No stop codon was identified, which indicates that this open frame is

not terminated. Hence the molecular mass of the corresponding protein cannot be calculated and compared to the apparent molecular mass of 28,000 Da of the FMN reductase, estimated on SDS-PAGE as described in

5 Example 5.6. On the other hand, the NH₂-terminal sequence of the protein identified in this way by analysis of the open reading frames of the sequence of 694-bp is identical to that NH₂-terminal sequence of the proteins SnaC purified as described in Example 5.

10 Similarly, the two internal protein sequences of the FMN reductase described in 5.6 occur in the protein deduced from the sequenced portion. This confirms that the gene isolated from cosmid pIBV4 does indeed correspond to the protein FMN reductase described in

15 Example 5.6, designated SnaC (SEQ ID no. 7).

A study of the DNA fragments of S.

pristinaespiralis strain SP92 carried by cosmids pIBV1 to pIBV2 demonstrated the presence of several genes involved in the biosynthesis of pristinamycins II and pristinamycins I. The snaA, snaB and samS genes code for enzymes participating in the biosynthesis of pristinamycins II, pristinamycin III synthase and probable SAM synthase, and are grouped together physically on a large DNA fragment cloned into plasmid pIBV1. Similarly, the snbA, snbR, papA and papM genes - which code for proteins participating in the biosynthesis of pristinamycins I, 3-hydroxypicolinic acid:AMP ligase (SnbA), the protein SnbR probably

responsible for the transport of pristinamycins I, the protein Papa involved in the biosynthesis of p-aminophenylalanine from chorisimic acid, and p-aminophenylalanine (phenyl-N)- methyltransferase

5 (PapM) - are grouped together on a large DNA fragment cloned into cosmid pIBV2. Similarly, the snaA and snaD genes on the one hand - which code for proteins participating in the biosynthesis of pristinamycins II, the protein SnaD probably being a peptide synthase -

10 and the snbC, snbD and snbE genes on the other hand - which code for the 3 peptide synthases SnbC, SnbD and SnbE involved in the formation of the peptide chain of pristinamycin I from its 6 separate amino acids - are grouped together on a large DNA fragment cloned into

15 cosmid pIVB3. These results confirm the hypothesis of the grouping together of the genes for the biosynthesis of pristinamycins II, and also of the genes for the biosynthesis of pristinamycins I, and afford the possibility of cloning the other genes involved in

20 these biosyntheses, by chromosome walking, upstream and downstream of the regions studied.

Furthermore, it is possible by hybridization of the total DNA of the different strains producing streptogramins (see Table 1) with the snaA, snaB, snaC, snaD, samS, snbA, snbR, snbC, snbD, snbE, papA and papM genes, or with the genes identified by chromosome walking, or with fragments of these genes, to isolate the genes corresponding to the same functions in the

other microorganisms producing streptogramins. This makes it possible, by the same approach as that envisaged for the pristinamycins, to isolate all the genes involved in the biosynthesis of the different 5 streptogramins.

EXAMPLE 9: Genetic study of DNA fragments by gene disruption

This example illustrates how it is possible to demonstrate the involvement of genes in the 10 biosynthesis of streptogramins by constructing strains derived from a producing strain and mutated in these genes by disruption, and by analysing the phenotype of such mutants. This example shows, furthermore, how to obtain strains that are left producing only one or 15 other of the A and B components of streptogramins, or producing A and B components with ratios different from those observed with the strain SP92.

9.1. Construction of a mutant of *S. pristinaespiralis* SP92 disrupted in the *snaA* gene

20 This example illustrates how it is possible, by disruption of the *snaA* gene, to construct a strain of *S. pristinaespiralis* SP92 which no longer produces pristinamycin IIA and which produces, in contrast, pristinamycin IIB.

25 This mutant was constructed for the purpose of confirming the functionality of the protein SnaA and of providing an intermediate of pristinamycin II production capable of being modified subsequently.

Its construction was carried out using a suicide vector capable of replicating in *E. coli* only but carrying a resistance marker which is expressed in *Streptomyces*. This vector, pDH5, was developed by

5 Hillemann et al. (1991).

9.1.1. Construction of plasmid pVRC505

This example illustrates how it is possible to construct a plasmid which does not replicate in *S. pristinaespiralis* SP92 and which may be used to

10 disrupt the snaA gene by single homologous recombination.

Plasmid pVRC505 was constructed to produce the SP92 chromosomal mutant disrupted in the snaA gene from plasmid pXL2045 described in Example 6.3.

15 The 6-kb BamHI fragment, cloned into pXL2045 (Figure 16), was cut with the restriction enzymes EcoRI and PstI. After separation of the fragments thereby generated by electrophoresis on 1 % agarose gel, a 0.7-kb fragment containing the 5' end of the snaA gene

20 was isolated and purified by Geneclean (Bio101, La Jolla, California).

100 ng of vector pDH5 linearized by an EcoRI/PstI double digestion were ligated with 100 ng of the 0.7-kb fragment, as described in Example 3.3. A

25 clone carrying the desired fragment was isolated after transformation of the strain TG1. The recombinant plasmid was designated pVRC505. Plasmid pVRC505 was prepared as described in Example 2.1. Its restriction

map is presented in Figure 25.

9.1.2. Isolation of the SP92 mutant disrupted
in the snaA gene by homologous recombination

This example illustrates how the mutant of
5 S. pristinaespiralis SP92 disrupted in the snaA gene
was constructed.

This mutant was isolated by transformation of
the strain SP92 with the suicide plasmid pVRC505.

The preparation of the protoplasts and their
10 transformation were carried out as described in
D. Hopwood et al. (1985).

The strain SP92 was cultured in YEME medium,
34 % sucrose, 5 mM MgCl₂, 0.25 % glycine for 40 hours at
30°C. The mycelium was converted to protoplasts in the
15 presence of lysozyme, and 5 x 1 µg of pVRC505 were used
for the transformation (by the method employing PEG) of
the protoplasts. After overnight regeneration of the
protoplasts on R2YE medium (D. Hopwood et al. 1985),
the recombinants were selected by overlaying 3 ml of
20 SNA medium (D. Hopwood et al. 1985) containing
2.6 µg/ml of thiostrepton.

Of the 5 transformations carried out, 3
thiostrepton-resistant clones were isolated. This gives
a recombinant efficiency of less than 1 per µg of DNA.
25 These recombinants result from integration by single
homologous recombination between the snaA gene carried
by the chromosome of the strain SP92 and the 0.7-kb
fragment of the suicide plasmid pVRC505. The small size

of the fragment inserted into pVRC505, 0.7-kb, explains the low recombination efficiency.

The spores of the recombinants were isolated by plating out and growth on R2YE medium supplemented 5 with 400 µg/ml of thiostrepton, and plated out again on the same medium to obtain isolated colonies.

In order to verify the position of integration of plasmid pVRC505, various Southern blots of the total DNA of several recombinant clones, which 10 was digested with the appropriate restriction enzymes, were produced and hybridized with the vector pDH5 and the 0.7-kb fragment, used successively as probes after labelling by random priming (Random Primed DNA labeling kit, Boehringer Mannheim, France) with [α -³²P]-dCTP, as 15 described in Maniatis *et al.* (1989). The hybridization results show the appearance in the genome of the recombinant clones of an additional EcoRI-PstI band, of the size of the vector pDH5 and which hybridizes with the latter, as well as of an additional EcoRI-EcoRI 20 band which hybridizes with both the 2 probes. One of these mutants was designated SP92::pVRC505. This mutant corresponds well to the integration of plasmid pVRC505 in the snaA gene by single homologous recombination.

9.1.3. Production of pristinamycins by the 25 mutant SP92::pVRC505

This example illustrates how it is determined that the mutant of S. pristinaespiralis SP92 disrupted in the snaA gene by integration of plasmid pVRC505 no

longer produces pristinamycin IIA while continuing to produce pristinamycin IIB.

The mutant SP92::pVRC505, as well as the strain SP92 as control strain, were culture in liquid 5 production medium. Fermentation was carried out as follows: 0.5 ml of a suspension of spores of the strains mentioned are added under sterile conditions to 40 ml of inoculum medium in a 300-ml Erlenmeyer. The inoculum medium consists of 10 g/l of corn steep, 10 15 g/l of sucrose, 10 g/l of $(\text{NH}_4)_2\text{SO}_4$, 1 g/l of K_2HPO_4 , 3 g/l of NaCl, 0.2 g/l of $\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$ and 1.25 g/l of CaCO_3 . The pH is adjusted to 6.9 with sodium hydroxide before the introduction of calcium carbonate. The Erlenmeyers are stirred for 44 hours at 27°C on a 15 rotary stirrer at a speed of 325 rpm. 2.5 ml of the above culture when 44 hours old are added under sterile conditions to 30 ml of production medium in a 300-ml Erlenmeyer. The production medium consists of 25 g/l of soya flour, 7.5 g/l of starch, 22.5 g/l of glucose, 20 3.5 g/l of feeding yeast, 0.5 g/l of zinc sulphate and 6 g/l of calcium carbonate. The pH is adjusted to 6.0 with hydrochloric acid before the introduction of calcium carbonate. The Erlenmeyers are stirred for 24, 28 and 32 hours at 27°C. At each time, 10 g of must are 25 weighed into a smooth Erlenmeyer, and 20 ml of mobile phase composed of 62.5 % of acetonitrile and 37.5 % of 0.1 M KH_2PO_4 solution (adjusted to pH 3.0 with H_3PO_4), and which enables the pristinamycins to be extracted,

are added to this. After stirring on a stirrer (325 rpm) for 20 min at room temperature, the whole is filtered through filter paper and the pristinamycins are assayed by HPLC as described in Example 5.1.1.A.

5 The results showed that, under the fermentation conditions implemented, the mutant SP92::pVRC505 produced an amount of pristinamycin I equivalent to that of the SP92 control, this being the case for all 3 times tested. In contrast, whereas the 10 control produced approximately 70 % of pristinamycin IIA and 30 % of pristinamycin IIB at 24, 28 and 32 hours of fermentation, the mutant SP92::pVRC505 produced 100 % of pristinamycin IIB for these same times, in amounts equivalent to the sum of 15 pristinamycin IIA + pristinamycin IIB produced by the strain SP92. Hence the mutant is indeed blocked in a step of biosynthesis of pristinamycin II which corresponds to the oxidation of the 2,3 bond of the D-proline of the intermediate pristinamycin IIB. This 20 mutant hence accumulates pristinamycin IIB. This shows well the functional involvement of SnaA in the conversion of pristinamycin IIB to pristinamycin IIA.

25 This example shows that it is possible, starting from cloned genes for biosynthesis, to construct strains that are mutated in the steps of biosynthesis of pristinamycin. This was shown for pristinamycin II, but the same results may be obtained for pristinamycins I and, by extension, for the

different components of streptogramins. Strains producing different intermediates may thus be obtained.

These strains may be used to produce novel molecules by chemical, biochemical, enzymatic, and the like,

5 modification(s) of the said intermediates. A block in an early step of the biosynthesis pathway of one or other of the components of streptogramins may also lead to mutated strains that are left producing only one or other of the components.

10 9.2. Construction of a mutant of *S.*

pristinaespiralis SP92 disrupted in the *samS* gene

This example illustrates how it is possible, by disruption of the *samS* gene, to construct a strain of *S. pristinaespiralis* SP92 which produces 35 % less PIA and 10 times as much PIB (the chemical structures are shown in Figure 2) relative to the wild-type SP92 strain. This mutant was constructed for the purpose of confirming the presumed SAM synthase function for the protein encoded by the *samS* gene, and for obtaining a

20 strain that synthesizes more PIB than the wild-type SP92 strain.

9.2.1. Construction of plasmid pVRC702

From plasmid pXL2045 (described in Example 6.3), the 3.2-kb BamH1-EcoR1 fragment was isolated by 25 enzymatic cleavage and purified after electrophoresis on 1 % agarose gel by the Geneclean kit method (see Example 6.8). This fragment carries snaB gene as well as the samS gene (Figure 16). This fragment is then

cloned into a plasmid pUC18 in the following manner:

50 ng of pUC18 were linearized by double digestion using the enzymes EcoR1 and BamH1, and then ligated in the presence of T_S DNA ligase (Biolabs) with 300 ng of 5 the 3.2-kb BamH1-EcoR1 fragment. After transformation of competent cells of E. coli strain TG1 with this ligation mixture, a recombinant clone possessing plasmid pUC18 with the 3.2-kb insert could be isolated, and this was designated pVRC701 (Figure 26).

10 Plasmid pVRC702 is derived from plasmid pVRC701 by the introduction between the two SstI sites located in the middle of the samS gene (Figure 27) of a cassette carrying the amR gene coding for resistance to amramycin and genitacin. To this end, a 2.2-kb BamH1-
15 BamH1 fragment carrying the QamR cassette was first isolated by BamH1 digestion of plasmid pHP450amR (given by J. L. Pernodet, Laboratoire de Génétique d'Orsay) using the same technique as above. 200 ng of this fragment were then ligated with 50 ng of plasmid
20 pUC1318 (Kay and McPherson, 1987) linearized with the enzyme BamH1, and this ligation mixture was introduced into competent E. coli TG1 cells. From the recombinant zone possessing plasmid pUC1318 containing the QamR cassette, 50 ng of a 2.2-kb SstI-SstI fragment
25 containing the QamR cassette could be reisolated by partial cleavage using the enzyme SstI, and this fragment was ligated with 30 ng of plasmid pVRC701 cut with SstI (Figure 26) to give, after transformation of

competent E. coli TG1 cells, plasmid PVRC702, the structure of which is detailed in Figure 27.

Plasmid PVRC702 thereby obtained was prepared in large amounts according to the method described 5 above in Example 2.1.

9.2.2. Construction of the strain having the samS::QamR chromosomal gene

This strain was obtained by transformation of S. pristinaespiralis protoplasts with 1 µg of the 10 suicide plasmid PVRC702 which is incapable of replicating in a Streptomyces cell. The protocols for preparation of the protoplasts and for transformation are the same as above (Example 9.1). The only 15 modifications made with respect to Example 10.1 relate to the selection antibiotic. In the present case, the recombinant protoplasts after regeneration for 18 hours at 30°C on R2YE medium are selected in the presence of 50 µg/ml final of geniticide (Sigma Chemical Co.). Thus, an overlayer composed of 3 ml of SNA containing 20 383 µg/ml of geniticide is added to each dish of R2YE.

In this way, it was possible to isolate 500 geniticide-resistant recombinant clones, which may result either from an integration of plasmid PVRC702 into the chromosome following a single homologous 25 recombination between chromosomal and plasmid samS genes (in the case of single crossing-over), or from an exchange between the chromosomal samS gene and the plasmid samS::QamR plasmid gene following a double

homologous recombination event (in the case of double crossing-over). In these two cases in point, the QamR cassette becomes transferred onto the chromosome of the strain, and endows it with an amR resistance which is 5 stable over generations.

The recombinant clones were isolated by plating out and growth on HT7 medium containing 50 µg/ml final of geneticin, and then analyzed by the colony hybridization technique. Hybridization of the 10 clones with a first probe obtained as described in Example 9.1 from the 2.7-kb BamH1-EcoRI fragment originating from pVRC702 and corresponding to pUC18, as well as with a second probe corresponding to the 2.2-kb BamH1 fragment carrying the QamR cassette, enable the 15 cases of single crossing-over (hybridizing with both probes) to be distinguished from the cases of double crossing-over (hybridizing only with the second probe). The 3 clones resulting from a double crossing-over thereby selected were purified by plating out and 20 growth on YVD medium containing 50 mg/ml final of geneticin, and stocks of spores were obtained.

In order to verify the genomic structure of the 3 double recombinants, various Southern blots of the chromosomal DNA of these clones digested with the 25 enzymes EcoRI and BamHI were produced and hybridized with the following three probes: the probe corresponding to the QamR cassette obtained from the 2.2-kb BamHI fragment of pVRC702, the probe

corresponding to pUC18 obtained from the 2.7-kb BamHI-EcoRI fragment of pVRC701, and lastly a probe obtained from the 1.3-kb EcoRI-SstI fragment of pVRC701 carrying the snaB gene and the beginning of samS. These 5 hybridizations enabled it to be verified that the three clones tested did indeed result from a double homologous recombination event permitting replacement of the intact chromosomal samS gene by the mutated sams gene interrupted by the QamR cassette.

10 One of these three mutant clones was designated SP92 samS::QamR.

9.2.3. Production of pristinamycins by the mutant strain samS::QamR

This example illustrates how it is determined 15 that the mutant SP92 samS::QamR having the disrupted samS gene produces 35 % less pristinamycin IA and 10-fold more pristinamycin IB than the wild-type SP92 strain.

The mutant SP92 samS::QamR as well as the 20 control SP92 strain were cultured in liquid production medium, and their productions of pristinamycin II and pristinamycin I were assayed as described in Example

9.1. —

The results showed that, under the 25 fermentation conditions implemented, the mutant SP92 samS::QamR produces an amount of pristinamycins II equivalent to that of the SP92 control for all three times tested. In contrast, the mutant SP92 samS::QamR

produces approximately 35 % less pristinamycin IA and 10-fold more pristinamycin IB than the control strain at all three times tested. The IB form of pristinamycins thus represents 20 % of the collective 5 total type I pristinamycins produced by the mutant SP92 samS::QamR, whereas the control strain synthesizes only of the order of 1 % of PIB. The IB form of pristinamycins differs from the IA form in that the fifth residue is p-methylaminophenylalanine, instead of 10 p-dimethylaminophenylalanine for pristinamycin IA. The fact that the mutant SP92 samS::QamR produces more pristinamycin IB and less pristinamycin IA shows that disruption of the samS gene causes a decrease in the degree of methylation of the fifth residue of 15 pristinamycins I, and hence that the samS gene is probably involved in the biosynthesis of the methyl donor, SAM, that is to say that it codes for a SAM synthase.

9.3 Construction of a mutant of S.

20 pristinaespiralis SP92 disrupted in the papA gene

This example illustrates how it is possible, by disruption of the papA gene, to construct a strain of S. pristinaespiralis SP92 which no longer produces PI. This mutant is constructed for the purpose of 25 confirming the functionality of the PapA protein. Its construction was carried out using the suicide vector pDH5 described in Example 9.1.

9.3.1. Construction of plasmid pVRC508

This example illustrates how it is possible to construct a plasmid which does not replicate in S. pristinaespiralis SP92 and which may be used to disrupt the papA gene by single homologous recombination.

Plasmid pVRC508 was constructed to produce the SP92 chromosomal mutant disrupted in the papA gene from plasmid pVRC903 described in Example 7.7.

In Example 7.7, the cloning of the 1.4-kb 10 PvuII-EcoRI fragment into M13mp18 from plasmid pVRC903 for the purpose of sequencing the papA gene was described (this fragment corresponds to the 1.4-kb PvuII-XhoI fragment present in the vector pVRC900, Figure 23).

15 This construction in M13mp18 was digested with the restriction enzyme HindIII and EcoRI. After separation of the vector M13mp18 and the 1.4-kb fragment containing a portion of the papA gene on 0.8 % agarose gel, the latter fragment was isolated and 20 purified by Geneclean (Biol01, La Jolla, California). The localization of the fragment in the papA gene is presented in Figure 23.

100 ng of vector pDH5 linearized by a double digestion with the restriction enzymes HindIII and 25 EcoRI were ligated with 200 ng of the 1.4-kb fragment as described in Example 3.3. A clone carrying the desired fragment was isolated after transformation of the strain TG1. The recombinant plasmid was designated

pVRC508. Plasmid pVRC508 was prepared as described in Example 2.1. Its restriction map is presented in Figure 28.

9.3.2. Isolation of the SP92 mutant disrupted
5 in the papA gene by homologous recombination

This example illustrates how the mutant of S. pristinaespiralis SP92 disrupted in the papA gene was constructed. This mutant was isolated by transformation of the strain SP92 with the suicide 10 plasmid pVRC508. The preparation of the protoplasts and their transformation were carried out as described in Example 9.1. After transformation of protoplasts of the strain SP92, the recombinants were selected by overlaying 3 ml of SNA medium containing 2.6 mg/ml of 15 thiostrepton. Of the 5 transformations carried out with 5 times 1 μ g of plasmid pVRC508, ten thiostrepton-resistant clones were isolated. This gives a recombinant efficiency of approximately 2 per μ g of DNA. These recombinants result from integration by 20 single homologous recombination between the papA gene carried by the chromosome of the strain SP92 and the 1.4-kb fragment of the suicide plasmid pVRC508.

The spores of the recombinants were isolated by plating out and growth on R2YE medium containing 25 400 μ g/ml of thiostrepton, and plated out again on the same medium to obtain isolated colonies.

In order to verify the position of integration of plasmid pVRC508, various Southern blots

of the total DNA of several recombinant clones, purified as described above, were produced and hybridized with the vector pDH5 and the 1.4-kb fragment, used successively as probes after labelling 5 by random priming with [α - 32 P]dCTP as described in Maniatis *et al.* (1989). The hybridization results show the disappearance from the genome of the recombinant clones digested with the restriction enzyme EcoRI (site flanking the 1.4-kb fragment) of the 6.8-kb EcoRI band, 10 and the appearance of two additional bands relative to the control SP92 strain, one of 2.4 kb hybridizing with the 1.4-kb fragment, and the other of 10.5 kb hybridizing both with pDH5 and with the 1.4-kb fragment. Digestion of the recombinant clones with the 15 restriction enzyme PstI shows the appearance of two additional bands relative to the control SP92 strain, one of 1.0 kb hybridizing with the 1.4-kb fragment, and the other of 5.1 kb hybridizing both with pDH5 and with the 1.4-kb fragment. One of these mutants was 20 designated SP92::pVRC508.

9.3.3. Production of pristinamycins by the mutant SP92::pVRC508

— This example illustrates how it is determined that the mutant of S. pristinaespiralis SP92 disrupted 25 in the papA gene by integration of plasmid pVRC508 no longer produces PI.

The mutant SP92::pVRC508, as well as the strain SP92 as control strain, were cultured in liquid

production medium. The fermentation and also the assay of pristinamycins I and II were carried out as described in Example 9.1.

The results showed that, under the 5 fermentation conditions implemented, whereas the control SP92 strain produced a standard amount of pristinamycins I, no trace of type I pristinamycins was detected in the fermentation must of the mutant SP92::pVRC508. Moreover, the production of 10 pristinamycins II by the mutant SP92::pVRC508 is equivalent to that of the SP92 control. The mutant SP92::pVRC508 is left producing only pristinamycins II. These results show clearly that the papA gene codes for a protein involved in the biosynthesis of 15 pristinamycins I.

To check the absence of polarity of the disruption carried out in the mutant SP92::pVRC508, the latter was fermented in the presence p-dimethylamino-phenylalanine. The mutant SP92::pVRC508 was fermented 20 as described above, with the addition, at 17 hours of fermentation, of 100 mg/l of p-dimethylaminophenylalanine. Under these conditions of complementation, the mutant SP92::pVRC508 produces an amount of pristinamycins I equivalent to that produced by the 25 strain SP92. The production of pristinamycins II is equivalent in both strains. This enables us to conclude that the mutant SP92::pVRC508 does not produce pristinamycins I because it is indeed disrupted in a

gene that participates in the biosynthesis of p-dimethylaminophenylalanine (the papA gene).

Complementation of this mutant with p-dimethylamino-phenylalanine restores its capacity to produce

5 pristinamycins I, proving that the mutation has no polar effect on the synthesis of other pristinamycin I precursors or on the condensation of these precursors.

This example shows that it is possible, starting from cloned genes for biosynthesis, to

10 construct strains that are mutated in the steps of biosynthesis of pristinamycins, and especially pristinamycins I. This example also shows that it is possible, by this approach, to construct strains of S. pristinaespiralis specifically producing

15 pristinamycins II and, by extension, strains specifically producing pristinamycins I. This same approach could also be used for other strains of actinomycetes producing streptogramins.

9.4. Construction of mutant of

20 S. pristinaespiralis SP92 disrupted in the snbA gene

This example illustrates how it is possible, by disruption of the snbA gene, to construct a strain of S. pristinaespiralis SP92 which no longer produces pristinamycins I. This mutant was constructed for the 25 purpose of confirming the functionality of the SnbA protein. Its construction was carried out using the suicide vector pDH5 described in Example 9.1.

9.4.1. Construction of plasmid pVRC404

This example illustrates how it is possible to construct a plasmid which no longer replicates in S. pristinaespiralis SP92 and which may be used to disrupt the snbA gene by single homologous 5 recombination.

Plasmid pVRC404 was constructed from plasmid pVRC402 described in Example 6.2, to produce the SP92 chromosomal mutant disrupted in the snbA gene. Plasmid pVRC402 was digested with the restriction enzyme XbaI and HindIII. After separation of the fragments thereby generated by electrophoresis on 0.8 % agarose gel, a 1170-bp fragment containing an internal portion of the snbA gene was isolated and purified by Geneclean 10 (Biol01, La Jolla, California). The localization of the 15 fragment in the snbA gene is presented in Figure 15A.

100 ng of vector pDH5 linearized by an SmaI digestion were ligated with 200 ng of the 1173-bp fragment as described in Example 3.3. A clone carrying the desired fragment was isolated after transformation 20 of the strain TG1. The recombinant plasmid was designated pVRC404. Plasmid pVRC404 was prepared as described in Example 2.1. Its restriction map is presented in Figure 29.

9.4.2. Isolation of the SP92 mutant disrupted 25 in the snbA gene by homologous recombination

This example illustrates how the mutant of S. pristinaespiralis SP92 disrupted in the snbA gene was constructed.

This mutant was isolated by transformation of the strain SP92 with the suicide plasmid pVRC404. The preparation of the protoplasts and their transformation were carried out as described in Example 9.1. After 5 transformation of protoplasts of the strain SP92, the recombinants were selected by overlaying 3 ml of SNA medium containing 2.6 mg/ml of thiostrepton. Of the 5 transformations carried out with 5 times 1 μ g of plasmid pVRC404, about thirty thiostrepton-resistant 10 clones were isolated. This gives a recombinant efficiency of approximately 5 per μ g DNA. These recombinants result from integration by single homologous recombination between the snbA gene carried by the chromosome of strain SP92 and the 1170-bp 15 fragment of the suicide plasmid pVRC404. The spores of the recombinants were isolated by plating out and growth on R2YE medium + 400 mg/ml of thiostrepton, and plated out again on the same medium to obtain isolated colonies. In order to verify the position of 20 integration of plasmid pVRC404, various Southern blots of the total DNA of several recombinant clones, purified as described above, were produced and hybridized with the vector pDH5 and the 1170-kb fragment, used successively as probes after labelling 25 by random priming with [α -³²P]dCTP as described in Maniatis et al. (1989). The hybridization results show the appearance in the genome of the recombinant clones digested with the restriction enzymes XhoI and HindIII

of an additional 4.7-kb XbaI-HindIII band relative to the control SP92 strain (vector pDH5 + 1.17 kb), hybridizing both with pDH5 and with the 1170-bp fragment. Digestion of the recombinant clones with the 5 restriction enzyme PflMI (sites flanking the 1170-bp XbaI-HindIII fragment) shows the disappearance of the 3.1-kb PflMI-PflMI band and the appearance of a band at 8.8 kb hybridizing with both probes. These results indicate that the genomic structure of the clones 10 analysed is indeed that expected after a homologous recombination event between pVRC404 and the chromosomal snbA gene. One of these mutants was designated SP92::pVRC404.

9.4.3. Production of pristinamycins by the 15 mutant SP92::VRC404

This example illustrates how it is determined that the mutant of S. pristinaespiralis SP92 disrupted in the snbA gene by integration of plasmid pVRC404 no longer produces PI.

20 The mutant SP92::pVRC404, as well as the strains SP92 as control strain, were cultured in liquid production medium. The fermentation and also the assay of pristinamycins I and II were carried out as described in Example 9.1. The results showed that, 25 under the fermentation conditions implemented, whereas the control SP92 strain produces a standard amount of pristinamycins I, no trace of pristinamycins I was detected in the fermentation must of the mutant

SP92::pVRC404. Moreover, the production of pristinamycins II by the mutant SP92::pVRC404 is equivalent to that of the SP92 control. The mutant SP92::pVRC404 is left producing only pristinamycins II.

5 This shows clearly that the snbA gene codes for a protein SnbA involved in the biosynthesis of pristinamycins I, as had been shown during the purification in Example 5.2.

This example shows, as in the preceding 10 example, that it is possible, starting from cloned genes for biosynthesis, to construct strains that are mutated in the steps of biosynthesis of pristinamycins, and especially pristinamycins I. This example also shows that is possible, by this approach, to produce 15 strains of S. pristinaespiralis specifically producing pristinamycins II and, by extension, strains specifically producing pristinamycins I, as described in the following example: 9.5. This same approach could also be used for other strains of actinomycetes 20 producing streptogramins.

9.5. Construction of a mutant of
S. pristinaespiralis SP92 disrupted in the snaD gene
probably coding for a peptide synthase involved in the
biosynthesis of pristinamycins II

25 This example illustrates how it is possible, by disruption of the snaD gene probably coding for a peptide synthase involved in the biosynthesis of pristinamycins II, to construct a strain of

S. pristinaespiralis SP92 which no longer produces pristinamycins II.

This mutant was constructed for the purpose of confirming the functionality of the snaD gene, and 5 of obtaining a strain derived from SP92 left synthesizing only pristinamycins I.

Its construction was carried out using plasmid PVRC1000 described in Example 6.8, derived from the suicide vector pDE5, capable of replicating in 10 E. coli only and carrying a resistance marker which is expressed in Streptomyces (see Example 9.1).

9.5.1 Construction of plasmid PVRC1000

This example illustrates how it is possible to construct a plasmid which does not replicate in 15 S. pristinaespiralis SP92 and which may be used to disrupt the snaD gene by single homologous recombination. The construction of plasmid PVRC1000 carrying a portion of the snaD gene is described in Example 6.8.

9.5.2. Isolation of the SP92 mutant disrupted in the snaD gene by homologous recombination

This example illustrates how the mutant of S. pristinaespiralis SP92 disrupted in the snaD gene was constructed. This mutant was isolated by 25 transformation of the strain SP92 with the suicide plasmid PVRC1000. The preparation of the protoplasts and their transformation were carried out as described in Example 9.1. Of the 5 transformations carried out

with 1 mg of pVRC1000, approximately 1500 thiostrepton-resistant clones were isolated. This gives a recombinant efficiency of approximately 375 per μ g of DNA. These recombinants result from integration by 5 single homologous recombination between the snaD gene carried by the chromosome of the strain SP92 and the 1.5-kb BamHI-SstI fragment of the suicide plasmid pVRC1000. About twenty recombinants were subcultured on R2YE medium containing 400 μ g/ml of thiostrepton, and 10 the spores of these recombinants were isolated by plating out again and growth on R2YE medium containing 400 μ g/ml of thiostrepton.

In order to verify the position of integration of plasmid pVRC1000, various Southern blots 15 of the total DNA of 7 recombinant clones, purified as described above, were produced and hybridized with the vector pDH5 and the 1.5-kb BamHI-SstI fragment contained in pVRC1000, used successively as probes after labelling with [α -³²P]dCTP as described in Example 20 9.1. The hybridization results show the appearance in the genome of the 7 recombinant clones of a 13.8-kb EcoRI band and an approximately 17-kb BglII band hybridizing with both probes, as well as a 3.7-kb EcoRI band hybridizing with the 1.2-kb BamHI-StsI probe. One 25 of these mutants was designated SP92::pVRC1000 and corresponds well to the integration of plasmid pVRC1000 in the snaD gene by single homologous recombination.

9.5.3. Production of pristinamycins by the mutant SP92::pVRC1000

This example illustrates how it is determined that the mutant of S. pristinaespiralis SP922 disrupted 5 in the snaD gene by integration of plasmid pVRC1000 no longer produces pristinamycins II, but only pristinamycins I. The mutant SP92::pVRC1000, as well as the control SP92 strain, were cultured in liquid production medium, and their productions of 10 pristinamycins II and I were assayed as described in Example 9.1.

The results showed that, under the fermentation conditions implemented and for all three times tested, the mutant SP92::pVRC1000 produces 0 mg/l 15 of pristinamycins II and an amount of pristinamycins I equivalent to that of the SP92 control. Hence this mutant is indeed blocked in a step of biosynthesis of pristinamycins II, which shows that the snaD gene codes for an enzyme involved in the biosynthesis of 20 pristinamycins II, and very probably for a peptide synthase.

This example shows, as in the preceding example, that it is possible, starting from cloned genes for biosynthesis, to construct strains that are 25 mutated in the steps of biosynthesis of pristinamycins, and especially pristinamycins II. This example also shows that it is possible, by this approach, to produce strains of S. pristinaespiralis specifically producing

pristinamycins I and, by extension, strains specifically producing pristinamycins II. This same approach could also be used for other strains of actinomycetes producing streptogramins.

5 EXAMPLE 10: Complementation of a non-producing mutant of the strain SP92

This example shows how it is possible to express genes for the biosynthesis of pristinamycins. This expression was implemented more especially for the 10 snaA and snaB genes carried by cosmid pIVB1 in a mutant strain derived from SP92: SP120. This mutant does not produce pristinamycin IIA. It accumulates the last intermediate of the biosynthesis pathway of pristinamycin II: pristinamycin IIB.

15 10.1 Cloning of the snaA and snaB genes into the shuttle vector pIJ903

This example illustrates how a subfragment of cosmid pIVB1 containing the snaA and snaB genes was cloned into a vector capable of replicating both in 20 E. coli and in Streptomyces.

The vector pIJ903 (Lydiate D. J. et al., 1985) is a low copy number (1 to 3 per cell) shuttle vector capable of replicating both in E. coli as a result of its origin of replication of pBR322, and in 25 Streptomyces as a result of its origin of replication of SCP2'. The ampicillin resistance gene permits selection in E. coli, and the thiostrepton resistance gene permits selection in Streptomyces.

Cosmid pIBV1 was digested with the restriction enzyme SstI. A large 7.6-kb DNA fragment carrying the snaA and snaB genes was isolated by electrophoresis on 0.8 % agarose gel and electroeluted.

5 500 ng of this fragment were ligated with 100 ng of the vector pUC1813 (Kay and McPherson, 1987) linearized with SstI. After transformation of E. coli strain DH5 α (supE44 ΔlacU169 (f80_λlacZAM15) hsdR17 recA1 endA1 gyrA96 thi-1 relA1), and selection of the transformants 10 on solid LB containing 150 μ g/ml of ampicillin and 20 μ g/ml of X-gal, a clone carrying the 7.6-kb fragment was isolated. The plasmid was designated pVRC506. A preparation of this recombinant plasmid was carried out as described in Example 2.1.

15 Cloning into the vector pIJ903 was carried out at the HindIII site. Plasmid pVRC506 was cut with HindIII, and 7.6-kb fragment carrying the snaA and snaB genes was isolated by electrophoresis on 0.8 % agarose gel and electroeluted. 500 ng of this fragment were 20 ligated with 500 ng of the vector pIJ903 linearized with HindIII. After transformation of E. coli strain DH5 α and selection of the transformants on solid LB containing 150 μ g/ml of ampicillin, a clone carrying the 7.6-kb fragment was isolated. The plasmid was 25 designated pVRC507. A preparation of this recombinant plasmid was carried out as described in Example 2.1. Its map is presented in Figure 30.

10.2. Expression of the snaA and snaB genes

in the mutant SP120

This example illustrates how it is possible to produce the proteins SnaA and SnaB in *S. pristinaespiralis* SP92 by introducing a plasmid carrying the corresponding structural genes into this strain. Expression of the snaA and snaB genes was carried out after transformation of the mutant strain SP120 with 500 ng of plasmid pVRC507. Transformation of the protoplasts of SP120 and selection of the 10 transformants with thiostrepton were carried out as described in Example 9.1.2.

Many transformants were obtained in this way, and 3 of them were chosen for the tests of production in a liquid medium. The strain SP120 carrying plasmid 15 pIJ903 was chosen as control. The fermentations and also the extraction of the biosynthesis products were carried out as described in Example 9.1.3.

The results showed that, under the fermentation conditions implemented, whereas the 20 control (SP120 carrying plasmid pIJ903) produced 100 % of P IIB and 0 % of P IIA at 24, 28 and 32 hours of fermentation, the 3 clones of the strain SP120 transformed with plasmid pVRC507 produced, for these same times, approximately 85 to 80 % of pristinamycin 25 IIB and 15 to 20 % of pristinamycin IIA, the sum of which is equivalent in amount to the pristinamycin IIB production of the control strain (SP120 carrying plasmid pIJ903). The clones carrying pVRC507 were

indeed partially complemented for the step of biosynthesis of pristinamycins II corresponding to the oxidation of the 2,3 bond of the D-proline of the intermediate pristinamycin IIB. This was confirmed by 5 enzymatic assay of pristinamycin IIA synthase activity, as described in Example 5.1.1.A, for the strains SP120 carrying pVRC507 and SP120 carrying pIJ903. Whereas the control strain SP120 carrying pIJ903 displays no pristinamycin IIA synthase activity, the strain SP120 10 carrying pVRC507 displays PIIA synthase activity.

This example shows that it is possible to express genes for the biosynthesis of streptogramins. This expression was studied more especially for the genes coding for pristinamycin IIA synthase, but the 15 other genes for the biosynthesis of pristinamycins II and pristinamycins I, as well as those involved in the biosynthesis of the components of the different streptogramins, may be expressed in this way. This expression may be carried out in mutant strains as is 20 the case in Example 10, but also in producing strains in order to increase the levels of streptogramin production. The expression may be modified by cloning the genes into a vector having a different copy number (low or high) or into an integrative vector, by 25 deregulation of these genes, by cloning these genes under a homologous or heterologous promoter (strong or specifically regulated promoter). Expression of the different genes for the biosynthesis of streptogramins

may also be carried out in heterologous strains using appropriate expression vectors in order to produce hybrid antibiotics.

EXAMPLE 11: Expression of the papM gene of
5 S. pristinaespiralis in E. coli

This example illustrates how it is possible to express an S. pristinaespiralis gene in E. coli so as to be able to identify, purify and study the protein encoded by this gene.

10 11.1. Construction of plasmid pVRC706

Expression of the papM gene in E. coli is obtained by placing this gene downstream of the promoter and ribosomes binding site of the lacZ gene of E. coli. The 1.7-kb MluI-StuI fragment was isolated
15 from plasmid pVRC409 described in Example 7.8, and then cloned into plasmid pMTL23 (Chambers et al. 1988) cut at the BamHI site subsequently filled in using the Klenow enzyme (Maniatis et al. 1989) and at the MluI site, to give plasmid pVRC706 shown in Figure 31.

20 Cloning at the MluI site enables an in-frame fusion to be obtained between the first 32 amino acids of β -galactosidase encoded by the lacZ gene of plasmid pMTL23 and the last eleven amino acids of the gene located immediately upstream of papM, thereby making it
25 possible to preserve the translational coupling which appears to exist between the papM gene and this upstream gene in the light of the nucleotide sequence given in Example 7.8. Thus, the expression of the

hybrid gene between lacZ and the gene upstream of pamM and that of the pamM gene is under the control of the expression signals of the lacZ gene.

5 11.2 Expression in *E. coli* strain DH5a of the product of the pamM gene

Plasmids pVRC706 and pMTL23 were introduced by transformation into *E. coli* strain DH5 α , and the expression of their genes was studied under conditions where the promoter of lacZ gene is induced as already 10 described (Maniatis *et al.* 1989). The *E. coli* strains carrying plasmid pVRC706 or the control plasmid pMTL23 were cultured in 500 ml of LB rich medium containing 100 mg/ml of ampicillin and 1 mM IPTG, permitting induction of the promoter of the lacZ gene. These 15 cultures are sampled when they have reached an optical density at 600 nm in the region of 1, and the protein extracts are prepared as described below.

20 11.3. Assay of the activity of the product of the pamM gene expressed in *E. coli*

The activity corresponding to the protein encoded by the pamM gene is assayed on the two extracts prepared from *E. coli* cultures carrying plasmid pVRC706 or plasmid pMTL23 (Example 5.7). It was shown that the extract prepared from the strain *E. coli*::pVRC706 25 catalyses the methylation of p-aminophenylalanine to p-dimethylaminophenylalanine with an activity of 235 unit/mg, whereas this activity is absent in the extracts of the control strain *E. coli*::pMTL23 (see

Example 5.7.1.C). These results indicate that it is possible to express the padM gene of S. pristinae-spiralis in E. coli, and that the corresponding protein is indeed the enzyme catalysing the methylation of p-aminophenylalanine to p-dimethylaminophenylalanine.

5 This example shows that it is possible to express genes for the biosynthesis of streptogramins in heterologous strains (such as E. coli, but also in other microorganisms) using appropriate expression vectors in

10 order to produce precursors of antibiotics or even natural or hybrid antibiotics.

EXAMPLE 12: Demonstration of the homology of genes involved in the biosynthesis of streptogramins in different Streptomyces

15 This example illustrates how it is possible to demonstrate, by hybridization with total DNAs, the homology existing between different genes involved in the biosynthesis of streptogramins in different strains of Streptomyces producing streptogramins.

20 **12.1. Extraction of total DNA of different Streptomyces producing streptogramins**

25 This example illustrates how the DNA of different strains producing streptogramins was purified. These strains of Streptomyces were chosen from those described in Table 1:

Streptomyces loidensis

Streptomyces olivaceus

Streptomyces ostreogriseusStreptomyces virginiae

A strain not producing streptogramins:

Streptomyces hygroscopicus, was chosen as negative

5 control.

The extractions of the different total DNAs were carried out from cultures in YEME medium, as described in Example 1.

10 12.2. Hybridization of total DNAs of strains producing streptogramins with DNA fragments containing genes involved in the biosynthesis of pristinamycins and isolated from S. pristinaespiralis strain SP92

15 This example illustrates how it is possible, starting from genes involved in the biosynthesis of pristinamycins and isolated from the strain SP92 as described in the preceding examples, to demonstrate homologous genes by hybridization of the total DNAs of strains producing streptogramins.

The DNA fragments used as a probe were:

20 The 3.9-kb XbaI-XbaI fragment isolated from pVRC1106 described in Example 6.5, the restriction map of which is presented in Figure 18. This fragment contains a portion of the gene coding for pristinamycin I synthase II.

25 The 6-kb BamHI-BamHI fragment isolated from plasmid pXL2045 described in Example 6.3, the map of which is presented in Figure 16. This fragment contains the structural genes for the two subunits of PIIA

synthase.

The total DNAs of the four strains producing streptogramins, the strain S. hygroscopicus and also the strain SP92, were digested with the restriction enzymes BamHI and XhoI. The DNA fragments thereby obtained were separated on 0.7 % agarose gel and the DNA was transferred onto a nylon membrane as described by Maniatis et al. (1989). Labelling of the 3.6-kb XhoI-XhoI and 6-kb BamHI-BamHI fragments was carried out by labelling by random priming as described in Example 9.1.2. Hybridization of the membranes was carried out in the presence of formamide at 42°C as described in Maniatis et al. (1989). Washing of the membranes after hybridization was carried out at 50 and 60°C in a solution containing SSC (Maniatis et al. (1989) diluted 10-fold and 0.1 % SDS.

The following results are demonstrated by these hybridizations:

The strain S. hygroscopicus does not display 20 and hybridization with the two probes used.

The total DNAs (digested with XhoI and BamHI) of the strains S. ostreogirseus, S. olivaceus, S. loidensis and S. virginiae all display hybridization signals of intensity comparable to those observed on 25 the total DNA of the strain SP92 with both probes used.

The total DNA (digested with XhoI and BamHI) of the strain S. virginiae displays signals with both probes used, but their intensity is weaker than that

observed in SP92.

This example shows that different strains of Streptomyces producing streptogramins contain genes that hybridize with genes isolated in

5 S. pristinaespiralis SP92 and which are involved in the biosynthesis of streptogramins, as presented in the preceding examples. These hybridizations thus demonstrate the homology existing between the genes involved in the biosynthesis of streptogramins of the
10 strains SP92 and those involved in the biosynthesis of streptogramins of other strains producing streptogramins.

This example hence shows that it is possible, starting from genes isolated from SP92 and involved in
15 the biosynthesis of streptogramins, to isolate by hybridization and cloning the homologous genes present in other strains producing streptogramins.

EXAMPLE 13: Study of the physical binding of the different S. pristinaespiralis SP 92 genes involved in the biosynthesis of pristinamycins I and pristinamycins II

5 This example illustrates how it is possible to study the physical binding of the S. pristinaespiralis SP 92 genes involved in the biosynthesis of pristinamycins I and II. This study was carried out for the purpose of showing that all these 10 genes are grouped together on the chromosome in a cluster, and that it is hence possible by chromosome walking from the genes already identified to isolate other genes involved in the biosynthesis of pristinamycins I and II. Such an approach may be 15 envisaged for the genes involved in the biosynthesis of other streptogramins.

13.1 Restriction enzymes used for pulsed-field electrophoresis

The S. pristinaespiralis SP92 genome is 20 composed of 70 % to 75 % of nucleotides containing the basis G and C. To cut its genome into a small number of large fragments, we used enzymes which recognize a sequence rich in AT, such as AseI (AT/TAAT) and SspI (AAT/ATT) but also HindIII (A/AGCTT), EcoRI (G/AATTC), 25 NdeI (CA/TATG) and ClaI (AT/CGAT).

13.2. S. pristinaespiralis strains used for pulsed-field electrophoresis

We used the chromosomal DNA of several

strains to study by pulsed-field electrophoresis the physical binding of the genes involved in the biosynthesis of pristinamycins I and pristinamycins II. We prepared inserts as described in Example 4.1 of the 5 chromosomal DNA of S. pristinaespiralis strain SP92, and also of the chromosomal DNA of the strains derived from SP92 whose construction is described in Examples 9.1 and 9.4. These are the strain SP92::pVRC505 in which the snaA gene has been disrupted by integration 10 of plasmid pVRC505 (Example 9.1), and the strain SP92::pVRC404 in which the snbA gene has been disrupted by integration of plasmid pVRC404 (Example 9.4). The latter two strains were included in this study since they enabled the snaA and snbB genes to be positioned 15 accurately on the chromosome map by exploiting the presence of sites which rarely cut chromosomal DNA, AseI, SspI, HindIII, EcoRI, NdeI and ClaI, in plasmids pVRC505 and pVRC404.

13.3. DNA probes used for hybridization of 20 the fragments isolated by pulsed-field electrophoresis

We used different DNA fragments to obtain radioactively labelled probes as is described in Example 9.1, which we hybridized with the fragments separated by pulsed-field electrophoresis after 25 enzymatic cleavage of the chromosomal DNA inserts of the three strains presented above. The probes are as follows: the 3,2-kb EcoRI-BamHI fragment isolated from plasmid pVRC701 carrying the snaB and samS genes (see

Example 9.2), the 1.5-kb BamH1-SstI fragment isolated from plasmid pVRC1000 carrying a portion of the snaD gene (see Example 6.8), the 1.1-kb XhoI-HindIII fragment isolated from plasmid pVRC402 carrying the 5 snbA gene (see Example 6.1), the 2.4-kb PstI-PstI fragment isolated from plasmid pVRC900 carrying papA gene (see Example 6.7) and the 1.5-kb XhoI-PstI fragment isolated from plasmid pVRC509 carrying the snaC gene (see Example 6.9).

10 13.4. Localization on the chromosome of the different genes involved in the biosynthesis of pristinamycins I and II and study of their physical binding

Hybridization of the chromosomal DNAs of 15 S. pristinaespiralis strains SP92, SP92::pVRC404 and SP92::pVRC505, cut by single digestions and double digestions using the six enzymes mentioned above, with the different probes described above lead to the general map shown in Figure 32: the position of major 20 sites has been indicated, together with the position and direction of transcription of the genes involved in the biosynthesis of pristinamycins PI and PII. Thus, it is possible to calculate the distance separating the 3 chromosomal regions containing the genes identified, 25 namely that of the snbA, snbR, papA and papM genes (cosmid pIBV2, Example 5.2), that of the snaA, snaB, samS, snaD, snbC, snbD and snbE genes (cosmids pIBV1 and 3, Example 5.1) and lastly that of the snaC gene

(cosmid pIBV4, Example 5.6). For example, the distance between the snaA and snbA genes has been evaluated at approximately 160-170 kb. This shows that the genes already identified are all contained in a region 5 covering only 200 kb of the chromosome of the S. pristinaespiralis strain, equivalent to less than 3 % of the total length of the genome, which we have been able to estimate at 7500 kb by the pulsed-field electrophoresis technique.

10 These results show that the genes involved in the biosynthesis of pristinamycins I and II are grouped together on the chromosome in a cluster, and that it is hence possible by chromosome walking from the genes already identified to isolate other genes involved in 15 the biosynthesis of pristinamycins I and pristinamycins II. More generally, it is possible, by chromosome walking from any gene involved in the biosynthesis of streptogramins, to identify the other genes involved in this biosynthesis.

TABLE 1

MICROORGANISMS	ANTIBIOTICS
FUNGI	
<u><i>Micromonospora</i></u> sp.	Vernamycins
STREPTOMYCES	
<u><i>S. albiorectus</i></u>	Virginiamycins
<u><i>S. conganesis</i></u> (ATCC13528)	F1370 A, B
<u><i>S. diastaticus</i></u>	Plauracins, Streptogramins
<u><i>S. graminofasciens</i></u>	Streptogramins
<u><i>S. griseus</i></u> (NRRL2426)	Viridogrisein (Etamycin)
<u><i>S. griseoviridus</i></u>	Griseoviridin
<u><i>S. griseoviridus</i></u> (FERMP3562)	Neoviridogriseins
<u><i>S. lavendulae</i></u>	Etamycins
<u><i>S. loidensis</i></u> (ATCC11415)	Vernamycins
<u><i>S. mitakaensis</i></u> (ATCC15297)	Mikamycins
<u><i>S. olivaceus</i></u> (ATCC12019)	Synergistins (PA114 A, B)
<u><i>S. ostreocriseus</i></u> (ATCC27455)	Ostreogrycins
<u><i>S. pristinaespiralis</i></u> (ATCC25486)	Pristinamycins
<u><i>S. virginiae</i></u> (ATCC13161)	Virgiuniamycins (Staphylocins)
ACTINOMYCETES	
<u><i>A. auranticolor</i></u> (ATCC31011)	Plauracins
<u><i>A. azureus</i></u> (ATCC31157)	Plauracins
<u><i>A. daghestanicus</i></u>	Etamycin
<u><i>A. philippinensis</i></u>	A-2315 A, B, C
<u><i>Actinoplanes</i></u> sp. (ATCC33002)	A15104
<u><i>Actinoplanes</i></u> sp.	A17002 A, B, C, F
<u><i>Actinomadura flava</i></u>	Madumycins

Abbreviations used:

DNA: deoxyribonucleic acid

AMP: adenosine 5'-monophosphate

ATP: adenosine 5'-triphosphate

5 ETB: ethidium bromide

bis-tris: (bis[2-hydroxyethyl]iminotris[hydroxymethyl]-methane)

bis-tris propane: (1,3-bis[tris(hydroxymethyl)-methylamino]propane)

10 BSA: bovine serum albumin

HPLC: high performance liquid chromatography

OD: optical density

DTE: dithioerythritol

DTT: dithiothreitol

15 E64: trans-epoxysuccinyl-L-leucylamido-(4-guanidino)butane

EDTA: ethylenediaminetetraacetic acid

EGTA: ethylene glycol bis(β-aminoethyl)tetraacetic acid

20 FMN: flavin mononucleotide

FMNH₂: reduced flavin mononucleotide

Hepes: (N-[2-hydroxyethyl]piperazine-N'-(2-ethanesulphonic acid])

IPTG: isopropyl β-D-thiogalactopyranoside

25 kDa: kilodalton

kb: kilobase

LB: Luria broth (rich growth medium for E. coli)

NAD: nicotinamide dinucleotide

NADH:	reduced nicotinamide dinucleotide
PAGE:	polyacrylamide gel electrophoresis
bp:	base pair
PMSF:	phenylmethylsulphonyl fluoride
5 PPi:	pyrophosphate
rpm:	revolutions per min
A.S.:	ammonium sulphate
SAM:	S-adenosylmethionine
SDS:	sodium dodecyl sulphate
10 STI:	soybean trypsin inhibitor
TE:	buffer comprising 10 mM Tris-HCl, 1 mM EDTA, pH 7.5
Tris:	2-amino-2-hydroxymethyl-1,3-propanediol
UV:	ultraviolet rays
15 Xgal:	5-bromo-4-chloro-3-indoyl β -D-galactoside
YEME:	yeast extract-malt extract medium (rich growth medium for <u>Streptomyces</u>)
PEG:	Polyethylene glycol
LMP:	Low melting point
20 MW:	Molecular weight

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SEQUENCE LISTING

(1) GENERAL INFORMATION:

(i) APPLICANT

5 (A) NAME: RHONE-POULENC RORER S.A.
(B) STREET: 20, avenue Raymond ARON
(C) CITY: ANTONY
(E) COUNTRY: FRANCE
(F) POST CODE: 92165

10 (ii) TITLE OF INVENTION: POLYPEPTIDES INVOLVED IN
THE BIOSYNTHESIS OF STREPTOGRAMINS, NUCLEOTIDE
SEQUENCES CODING FOR THESE POLYPEPTIDES AND USES.

(iii) NUMBER OF SEQUENCES: 16

(iv) COMPUTER READABLE FORM:

15 (A) MEDIUM TYPE: Tape
(B) COMPUTER: IBM PC compatible
(C) OPERATING SYSTEM: PC-DOS/MS-DOS
(D) SOFTWARE: PatentIn Release #1.0,
Version #1.25 (EPO)

(2) INFORMATION FOR SEQ ID NO: 1:

20 (i) SEQUENCE CHARACTERISTICS

(A) LENGTH: 5392 base pairs
(B) TYPE: nucleic acid

(C) STRANDEDNESS: double

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

5 (iii) ANTI-SENSE: NO

(vi) ORIGINAL SOURCE

(A) ORGANISM: *Streptomyces**pristinaespiralis*

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:

GGATCTGGC	GTCCGCCGTC	AAGAACTGAA	CCGAGGGAGAC	ACCCACCATG	ACCGCACCCC	60
GCGGGCGCAT	CACCCCTGCC	GGCATCATCG	ACGGCCCCGG	CGGCCATGTG	GCCGCCCTGGC	120
GCCACCCGGC	GACCAAGGGCG	GACGCCAGC	TGACTTCGA	ATTCACCCGC	GACAAACGCC	180
GCACCCCTCGA	ACGCGGGCTG	TTGACGCCG	TGTTCATCGC	GGACATGTC	GCCGTGTGGG	240
GCACCCGGCT	GGACTCCCTG	TGCCGCACCT	CGCGCACCGA	GCACTTCGAA	CCGCTCACCC	300
TGCTCGCCGC	CTACGCCGCG	GTCACCGAGC	ACATCGGCCT	GTGCGCCACC	GCCACCAACCA	360
CGTACAACGA	ACCGGGCGCAC	ATCGCCGCC	CCTTCGCC	CCTCGACCAAC	CTCAGCGGCG	420
GCGGGGCCGG	CTGGAACGTC	GTCACCTCCG	CCGCACCGTG	GGAGTCCGCC	AACTTCGGCT	480
TCCCCGAGCA	CCTGGAGCAC	GGCAAAACGCT	ACGAGCGGGC	CGAGGGAGTTC	ATCGACGTCG	540

TCAAAAAACT GTGGGACAGC GACGGCCGCC CCGTCGACCA CCGCGGCACC CACITCGAGG 600
 CCCCCGGCCC GCTCGGGATC GCCCGCCCCC CGCAGGGCCG CCCCCGTCAATC ATCCAGGCCG 660
 GCTCTCTGGCC GGTGGGACGC GACTTCGCGG CCCCACCGC CGAGGTCAATC TTCAACCGGC 720
 ACAACCGGCT CTCCGACGCC CAGGACTTCT ACAGGCACCT CAAGGCACGC GTCGCCCGGC 780
 ACGGCCGCCGA CCCCAGAAG GTCCCTCGTGT GCGCGACCCCT CGCGCCGATC GTCGCCGCCA 840
 CCGACACCGA GGCGAAGCAG CGCTGCAGG AACTGCAGGA CCTCACCCAC GACCATGTCG 900
 CCTCTGCGAC CCTTCAGGAC CACCTCGCCG ACCTCGACCT GAGCGCGTAC CCGATCGACG 960
 GGGCCGTCCC CGACATCCCG TACACCAACC AGTCCCAGTC GACGACCGAG CGGCTGATCG 1020
 GCCTGGCCAG CGCGAGAAC CTCAACATCC GCGAGCTGGC CCTGCGGCTG ATGGGGACA 1080
 TCGTCTCGG CACACCGGAG CAGCTCGCCG ACCACATGGA GAGCTGGTTC ACCGGCCCGCG 1140
 GCGCCGACGG CTCAACATC GACITCCCCT ACCTGCCGGG CTCCGCCGAC GACTTCGTCG 1200
 ACCACGTGGT GCCCAGACTG CAGCGCCGCG GCCTGTACCG CTGGGCTAC GAGGGCACCA 1260
 CCTCTGCGGGC CAACCTCGGC ATCGACGCC CCGGAAAGCC AGGTGCAGCG GCTTGACTTC 1320
 CGTCCTAAAG GCGGGGGATT CCAGCGGTG CCGCGTGGG TTCCCTGCTTC ACCGACGACC 1380
 GCCCCGTCCG GGAGGACTCC CGTTGAGGTC TTATACCGTC TCCACAGGCC GAGGCCGCCA 1440
 GCGCCGCGGC CAGGATGTTG CGTGCCTGAT TCACTGCGCG GTCATGCACA GCGCCGCGAT 1500
 CGCACGTCCA CTCCCCGAGC TTCAAGGGCA CCTTCCCCCG GACCGTCCCG CAGGTTCCGC 1560
 ACAGCTTGGA CCTGGGGAAC CAGCGCTCGA TCACGACCGAG TTCCGCCCCA TACCAAGGCC 1620
 ACTTGTACTC CAGCATGGAG CGCAGTTCCG TCCAGGCCGC GTGGAGATG GCGCGCGCGA 1680
 GCTTGGCGTT TTTCAGCAGG TTGCGGACGG TGAGGTCTTC GATCACGACC GTTGGTTCT 1740
 CACGGACGAG TCGAGTCGAC AGCTTGTGGA GGAAGTCGCA GCGCCGGTGC GTGATCCGGG 1800
 CGTGGACCGG GGCGACCTTG CGGGGGCTT TCTTCCGGTT CGCCGACCCCC TTCCCTTGC 1860
 GCGACACGTC CCGCTGAGGC TTGGCGAGGC GGGCGCGGTC ACGGCGCTCG TGCTTGGGT 1920
 TGGTGTCTT CTCCCCGGTG GACAGGGTCA CCAGGGAGGT GATCCCCCGG TCGATCCCCA 1980
 CGGCCGCCGT GGTGGCGGGC GCGGGGGTGA TGGTGTCTTC GCACAGCAGG GACACGAACC 2040
 ACCGGCCCGC ACGGTCCGGG GACACGGTCA CGCTCGTGG CTCCGCCCT TCGGGAAAGGG 2100
GACGGGACCA GCGGATGTCC AGGGGCTCCG CGGTCTTCGC CAGCGTGAGC TGTCCGTAC 2160
 GCCACGTGAA GGCGCTGCCG GTGTACTCGG CCGACGGCTCT GGACTTTTTC CGCGACTTGT 2220
 ACCGGGGTA CTTCGACCGC TTGGGAAGA AGTGGCGAA CGCCGCTCGC AAGTGCAGCA 2280
 GCGCCCTGCTG GAGCGGGACG GAGGACACCT CGGAGACGAA GGCGACTTCT TCGGTCTTCT 2340
 TCCACTCCGT CAGCGCGGGC GACGACTGCA CGTAGGGAGAC CGGGCGCTGC TCGCCGTACC 2400

AGGCTCGCGT GCGCCCCCTCA AGCCCTTGT TGTACACGAG GCGGACACAG CCGAACGTGC 2460
 GGGACAGCTC AGCCGCCCTGC TCGTCCGTGG GATAAAAGCG GTACTTGAAA GCCCCTTGA 2520
 CCTGCTGCAT CACGCCCTCAC AGCCTATCAG TTCCCCGTGTG AGCGGGGGGT GTCTGCCGGT 2580
 GGTTCAGAC GCGAACCGC CCTGGCGGGC ATTGCCCAT CCTGCCCTG CTCCGCAAGA 2640
 GCTTCGTCTC CTCCCCGGTC TGAAGGCCGG GGTATCCACG AAGGAATTCT GATGACCGGG 2700
 CCCATCCCTCG TCGCCACCCCT CGACACCCGC GGCCCCGCCG CCACCCCTCGG CACGATCACC 2760
 CGCGCCGTGC GGGCCGCCGGA GGGCCGCCGGA TTCGACGCCGG TCCTGATCGA CGACCGGGCC 2820
 GCCGCCGGCG TCCAGGGCCG GTTCGAGACG ACGACGCTGA CGGCCCGCGT GGCCGCCGTC 2880
 ACCGAGCACA TCGGCCCTGAT CACCGCCCCCG CTCCCCGGCG ACCAGGGCCCC CTACCACTG 2940
 TCCCGGATCA CGCCCTCGCT CGACCCACCTC GCCCACGGCC GCACCGGGCTG GCTCGCGAGC 3000
 ACGGACACCA CGAACCCCGA GGGCCGCACC GGCGAACTCA TCGACGTCGT CGCGCCCTG 3060
 TGGGACAGCT TCGACGACGA CGCCCTCGTC CACGACCGCG CCGACGGCCG GTACTGGCGG 3120
 CTGCCCGCCG TCCACCAACT CGACCAACAG CGCAGGGACT TCGACGTCGC CGGGCCCCCTC 3180
 AACGTCGCCC GCCCCGCCGA GGGCCACCCCC GTCGTCGCCG TCACCGGCCG CGCCCTCGCC 3240
 CGCGCCGGCG ACCTCGCTT GCTCGACGAG CGGGCCGACG CGCCCTCGGT GAAGCAGCAG 3300
 GCACCGCAGC CCAAGATCCT CCTGCCCTG CCCCCCCCCGG CCCCCGAACI CCCCCGCCGAC 3360
 AGCCCCGGCG ACGGCTTCAC GTGGCGCGTC ACCGGCTCCG ACGACCCGGT CCTGGCGCG 3420
 CTCCCCCCCC GGGCCGGCCG CCGGGACCCG ACCGGGCCA CCACCCCTCGG CGAACGCCCTG 3480
 GGCTGGCCC GCCCCGAGAG CGGCCACGCC CTCACCAACCG CCTGACGACC CGTCCGGCCG 3540
 CTGCTTCTG GAGACTCATG TCCCGTCGCC TGTTCACCTC GGAGTCCGTG ACCGAGGGCC 3600
 ACCCCGACAA GATCGCCGAC CAGATCAGTG ACACCGTCT CGACGCCCTG CTGCGCGAGG 3660
 ACCCGCCCTC ACGCGTCGGC GTCGAGACCC TGATCACCAC CGGCCAGGTC CACATGGCG 3720
 GCGAGGTCAC CACCAAGGCG TACCGGCCA TCGCCCAACT GTCCCGCGAC ACGATCCCTG 3780
 CCATCGGCTA CGACTCGTCC CGCAAGGGCT TCGACGGCGC CCTCTGCCGGC GTCTCCGTCT 3840
 CCATCGGCGC GCAGTCCCCG GACATCGGCC AGGGCGTCGA CAGCGCTAC GAGACCCGGC 3900
 TCGAGGGCGA GGACGACCGAG CTGGACCCAGC AGGGCGCCGG CGACCCAGGGC CTGATGTCTG 3960
 GCTACGCCAT CGACGAGACC CCCTCGCTGA TGCCCGTCGCC CATCGAGCTC GCCCCACGCC 4020
 TCTCGCGCCG GCTCACCGAG GTCCGCAAGG ACGGCACCGT CCTCTACCTG CGCCCCGACG 4080
 GCAAGACCCA GGTCAACCATC GAGTACCAAGG GCAGCCGCCG GGTCCGCCCTG GACACCGTCG 4140
 TCGTCCTCTC CGAGCACGCC CGCGACATCG ACCTCGGCCCTC CCTGCTCACCC CCGACATCC 4200
 CGGAGCACGT CGTCGAGCAC GTCCCGCCG CACTCGCCGA GGACGCCATC AAGCTCGAGA 4260
 CGGACAACTA CGGCCCTCGTG GTCAACCCGA CGGGCCGTT CGAGATCGGC GGCCCGATGG 4320

GGGACGCCGG	CCTGACCGGC	CGCAAGATCA	TCATCGACAC	GTACGGCGGC	ATGGCCGCC	4380
ACGGCGGTGG	CGCGTTCTCC	GGCAAGGACC	CGTCCAAGGT	CGACCGTCC	GGCGCGTACG	4440
CGATGCCCTG	GGTCGCCAAG	AACGTCGTG	CCGCGGGCCT	CCCTCCCGC	TGGAGGTCC	4500
AGGTCGCCTA	CGCCATCGGC	AAGGCCGAGC	CGGTGCGCCT	GTTGTCGAG	ACGTTGGCA	4560
CCGGCACCGT	CGCCCAGGAG	CGCATCGAGA	AGGCCATCAC	CGAGGTCTTC	GACCTGGCC	4620
CCGCGGCCAT	CATCCCGGAC	CTCGACCTGC	TGCGGCCCAT	CTACGCCGCC	ACCGCCGCCT	4680
ACGGCCACTT	CGGCCGCGAA	CTGCCCAGT	TCACCTGGGA	GCGGACCGAC	CGCGCCACC	4740
GGCTCAAGGC	CGCGGCCGGT	CTCTGAGCCG	GCCGGACCTG	TGAGGAGACC	TGACGTGCGC	4800
ATCGCTGTCA	CCGGTTCCAT	CGCCACCGAC	CATCTGATGG	TCTTCCCCGG	CCGGTTCCCG	4860
GATCAGCTGA	TCCCCGACCA	GCTCGCTCAT	GTCTGGCTCT	CCTTCTGGT	CGACGCACTC	4920
GAGGTGCGCC	GGGGCGGAGT	GGCGGACAAC	GTCGCCCTCG	GCCTCGGCCG	CCTCGGCCCTC	4980
ACCCCCCAGC	TGGTCGGCGC	CGTGGGCAGC	GACTTCGCCG	AGTACGAGGT	CTGGCTCAAG	5040
GAACACGGCG	TCGACACCGG	CCCCGTCTG	GTCTCCACCG	AGCGGCAGAC	CCCCGGTTC	5100
ATGTGCATCA	CCGACCAGGA	CTCCAACCAG	ATCGCCTCCT	TCTACGGGG	CGCCATGCAA	5160
GAGCCCCCG	ACATCGACCT	GTGGCACCTG	ACCACCGGCA	GCGTCCGCC	CGACCTCGTC	5220
CTGGTCTGCC	CGAACGACCC	GGCGGCCATG	CTGCGCCACA	CGGGGAGTGC	CGCGAAACTG	5280
GGCTTCCCCGT	TGGCTGGCGA	CCCCCTCCAG	CAGCTCGCC	GCCTGGAGGG	AGGGAGGTAC	5340
CGGAACCTCGG	TCGACGGGGC	CCGTTGGTTT	TTCAACCAACG	AACTACGAGG	CC	5392

(2) INFORMATION FOR SEQ ID NO: 2:

(i) SEQUENCE CHARACTERISTICS

(A) LENGTH: 1269 base pairs

(B) TYPE: nucleic acid

5: (C) STRANDEDNESS: double

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iii) ANTI-SENSE: NO

10 (vi) ORIGINAL SOURCE

(A) ORGANISM: *Streptomyces*
pristinaespiralis

(ix) FEATURE:

(A) NAME/KEY: CDS

5 (B) LOCATION: 1..1269

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:

ATG ACC GCA CCC CGC CGG CGC ATC ACC CTC GCC GGC ATC ATC GAC GGC

Met Thr Ala Pro Arg Arg Arg Ile Thr Leu Ala Gly Ile Ile Asp Gly	15
1 5 10 15	
CCC GGC GGC CAT GTG GCC GCC TGG CGC CAC CCG GCG ACC AAG GCG GAC	96
Pro Gly Gly His Val Ala Ala Trp Arg His Pro Ala Thr Lys Ala Asp	
20 25 30	
GCC CAG CTC GAC TTC GAA TTC CAC CGC GAC AAC GCC CGC ACC CTC GAA	144
Ala Gln Leu Asp Phe Glu Phe His Arg Asp Asn Ala Arg Thr Leu Glu	
35 40 45	
GCG GGC CTG TTC GAC GCC GTG TTC ATC GCG GAC ATC GTC GCC GTG TGG	192
Arg Gly Leu Phe Asp Ala Val Phe Ile Ala Asp Ile Val Ala Val Trp	
50 55 60	
GCG ACC CGC CTG GAC TCC CTG TGC CGC ACC TCG CGC ACC GAG CAC TTC	240
Gly Thr Arg Leu Asp Ser Leu Cys Arg Thr Ser Arg Thr Glu His Phe	
65 70 75 80	
GAA CCG CTC ACC CTG CTC GCC TAC GCC GCG GTC ACC GAG CAC ATC	288
Glu Pro Leu Thr Leu Leu Ala Ala Tyr Ala Ala Val Thr Glu His Ile	
85 90 95	
GCG CTG TGC GCC ACC GCC ACC ACC ACG TAC AAC GAA CCG GCG CAC ATC	336
Gly Leu Cys Ala Thr Ala Thr Thr Tyr Asn Glu Pro Ala His Ile	
100 105 110	
GCC GCC CGC TTC GCC TCC CTC GAC CAC CTC AGC GGC GGC CGG CCC GGC	384
Ala Ala Arg Phe Ala Ser Leu Asp His Leu Ser Gly Gly Arg Ala Gly	
115 120 125	
TGG AAC GTC GTC ACC TCC GCC GCA CGG TGG GAG TCC GCC AAC TTC CGC	432
Trp Asn Val Val Thr Ser Ala Ala Pro Trp Glu Ser Ala Asn Phe Gly	
130 135 140	
TTC CCC GAG CAC CTG GAG CAC GGC AAA CCC TAC GAG CGG GCC GAG GAG	480
Phe Pro Glu His Leu Glu His Gly Lys Arg Tyr Glu Arg Ala Glu Glu	
145 150 155 160	
TTC ATC GAC GTC GTC AAA AAA CTG TGG GAC ACC GAC CCC CCC GTC	528
Phe Ile Asp Val Val Lys Lys Leu Trp Asp Ser Asp Gly Arg Pro Val	
165 170 175	
GAC CAC CGC GGC ACC CAC TTC GAG GCC CCC GGC CCG CTC GGG ATC GCC	576
Asp His Arg Gly Thr His Phe Glu Ala Pro Gly Pro Leu Gly Ile Ala	
180 185 190	
GCG CCC CCG CAG GGC CGC CCC GTC ATC ATC CAG GCC GGC TCC TCG CCG	624
Arg Pro Pro Gln Gly Arg Pro Val Ile Ile Gln Ala Gly Ser Ser Pro	
195 200 205	
GTC GGA GGG GAG TTC GCC GCC CGG CAC GCC GAG GTC ATC TTC ACC CGG	672
Val Gly Arg Glu Phe Ala Ala Arg His Ala Glu Val Ile Phe Thr Arg	
210 215 220	
CAC AAC CGG CTC TCC GAC GCC CAG GAC TTC TAC GGC GAC CTC AAG GCA	720
His Asn Arg Leu Ser Asp Ala Gln Asp Phe Tyr Gly Asp Leu Lys Ala	
225 230 235 240	
GCG GTC GCC CGG CAC GGC CGC GAC CCC GAG AAG GTC CTC GTG TGG CCG	768
Arg Val Ala Arg His Gly Arg Asp Pro Glu Lys Val Leu Val Trp Pro	
245 250 255	

ACC CTC GCG CCG ATC GTC GCC GCC ACC GAC ACC GAG GCG AAG CAG CGC	816
Thr Leu Ala Pro Ile Val Ala Ala Thr Asp Thr Glu Ala Lys Gln Arg	
260 265 270	
CTG CAG GAA CTG CAG GAC CTC ACC CAC GAC CAT GTC GCC CTG CGC ACC	864
Leu Gln Glu Leu Gln Asp Leu Thr His Asp His Val Ala Leu Arg Thr	
275 280 285	
CTT CAG GAC CAC CTC GGC GAC GTC GAC CTG AGC GCG TAC CCG ATC GAC	912
Leu Gln Asp His Leu Gly Asp Val Asp Leu Ser Ala Tyr Pro Ile Asp	
290 295 300	
GGG CCC GTC CCC GAC ATC CCG TAC ACC AAC CAG TCC CAG TCG ACG ACC	960
Gly Pro Val Pro Asp Ile Pro Tyr Thr Asn Gln Ser Gln Ser Thr Thr	
305 310 315 320	
GAG CGG CTG ATC GGC CTG GCC AGG CGC GAG AAC CTC AGC ATC CGC GAG	1008
Glu Arg Leu Ile Gly Leu Ala Arg Arg Glu Asn Leu Ser Ile Arg Glu	
325 330 335	
CTG GCC CTG CGG CTG ATG GGC GAC ATC GTC GTC GGC ACA CCG GAG CAG	1056
Leu Ala Leu Arg Leu Met Gly Asp Ile Val Val Gly Thr Pro Glu Gln	
340 345 350	
CTC GCC GAC CAC ATG GAG AGC TGG TTC ACC GGC CGC GGC GCC GAC GGC	1104
Leu Ala Asp His Met Glu Ser Tyr Phe Thr Gly Arg Gly Ala Asp Gly	
355 360 365	
TTC AAC ATC GAC TTC CCG TAC CTG CCG GGC TCC GCC GAC GAC TTC GTC	1152
Phe Asn Ile Asp Phe Pro Tyr Leu Pro Gly Ser Ala Asp Asp, Phe Val	
370 375 380	
GAC CAC GTG GTG CCC GAA CTG CAG CGC CGC GGC CTG TAC CGC TCG GGC	1200
Asp His Val Val Pro Glu Leu Gln Arg Arg Gly Leu Tyr Arg Ser Gly	
385 390 395 400	
TAC GAG GGC ACC ACC CTG CGG GCC AAC CTC GGC ATC GAC GCC CCC CGG	1248
Tyr Glu Gly Thr Thr Leu Arg Ala Asn Leu Gly Ile Asp Ala Pro Arg	
405 410 415	
AAG GCA GGT GCA GCG GCT TG	1269
Lys Ala Gly Ala Ala Ala	
420	

(2) INFORMATION FOR SEQ ID NO: 3:

(i) SEQUENCE CHARACTERISTICS

(A) LENGTH: 834 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: double

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iii) ANTI-SENSE: NO

(iv) ORIGINAL SOURCE

(A) ORGANISM: *Streptomyces*
pristinaespiralis

(ix) FEATURE:

(A) NAME/KEY: CDS

5

(B) LOCATION: 1..834

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:

ATG ACC GCG CCC ATC CTC GTC GCC ACC CTC GAC ACC CGC GGC CCC CCC	48
Met Thr Ala Pro Ile Leu Val Ala Thr Leu Asp Thr Arg Gly Pro Ala	
1 5 10 15	
GCC ACC CTC GGC ACG ATC ACC CGC GCG GTC CGG CCC GCG GAG CCC GCG	96
Ala Thr Leu Gly Thr Ile Thr Arg Ala Val Arg Ala Ala Glu Ala Ala	
20 25 30	
GGA TTC GAC GCC GTC CTG ATC GAC GAC CGG CCC CCC GGC GTC CAG	144
Gly Phe Asp Ala Val Leu Ile Asp Asp Arg Ala Ala Ala Gly Val Gln	
35 40 45	
GCC CGG TTC GAG ACG ACG ACC GTC ACC GCC GCG CTG CCC GCC GTC ACC	192
Gly Arg Phe Glu Thr Thr Leu Thr Ala Ala Leu Ala Ala Val Thr	
50 55 60	
GAG CAC ATC GGC CTG ATC ACC CCC CCG CTC CCG GCC GAC CAG GCC CCC	240
Glu His Ile Gly Leu Ile Thr Ala Pro Leu Pro Ala Asp Gln Ala Pro	
65 70 75 80	
TAC CAC GTG TCC CGG ATC ACC CCC TCG CTC GAC CAC CTC CCC CAC GGC	288
Tyr His Val Ser Arg Ile Thr Ala Ser Leu Asp His Leu Ala His Gly	
85 90 95	
CGC ACC GGC TCG CTC GCG AGC ACG GAC ACC ACC GAC CCC GAG GGC CGC	336
Arg Thr Gly Trp Leu Ala Ser Thr Asp Thr Asp Pro Glu Gly Arg	
100 105 110	
ACC GGC GAA CTC ATC GAC GTC GTC CGC CGC CTG TGG GAC AGC TTC GAC	384
Thr Gly Glu Leu Ile Asp Val Val Arg Gly Leu Trp Asp Ser Phe Asp	
115 120 125	
GAC GAC GCC TTC GTC CAC GAC CGC GCC GAC GGC CTG TAC TGG CGG CTG	432
Asp Asp Ala Phe Val His Asp Arg Ala Asp Gly Leu Tyr Trp Arg Leu	
130 135 140	
CCC GCC GTC CAC CAA CTC GAC CAC CAG GGC AGG CAC TTC GAC GTG GCC	480
Pro Ala Val His Gln Leu Asp His Gln Gly Arg His Phe Asp Val Ala	
145 150 155 160	
GGC CCC CTC AAC GTC GCC CGC CCG CCG CAG GGC CAC CCC GTC GTC GCC	528
Gly Pro Leu Asn Val Ala Arg Pro Pro Gln Gly His Pro Val Val Ala	
165 170 175	
GTC ACC GGC CCC GCC CTC GCG GCC GAC CTC GTC CTG CTC GAC	576
Val Thr Gly Pro Ala Leu Ala Ala Ala Asp Leu Val Leu Leu Asp	
180 185 190	

GAG GCG GCC GAC GCC TCG GTG AAG CAG CAG GCA CCG CAC GCC AAG 624
 Glu Ala Ala Asp Ala Ala Ser Val Lys Gln Gln Ala Pro His Ala Lys
 195 200 205
 ATC CTC CTG CCG CTG CCC GGC CCG GCC GAA CTG CCC GCC GAC AGC 672
 Ile Leu Leu Pro Leu Pro Gly Pro Ala Ala Glu Leu Pro Ala Asp Ser
 210 215 220
 CCC GCG GAC GGC TTC ACG GTG GCG CTC ACC GGC TCC GAC GAC CCG GTC 720
 Pro Ala Asp Gly Phe Thr Val Ala Leu Thr Gly Ser Asp Asp Pro Val
 225 230 235 240
 CTG GCC GCG CTC GCC GCC CGG CCC GGC CGC CGG GAC CGC ACC GCG GCC 768
 Leu Ala Ala Leu Ala Ala Arg Pro Gly Arg Pro Asp Arg Thr Ala Ala
 245 250 255
 ACC ACC CTG CGC GAA CGC CTG GGC CTG GCC CGC CCC GAG AGC CGC CAC 816
 Thr Thr Leu Arg Glu Arg Leu Gly Leu Ala Arg Pro Glu Ser Arg His
 260 265 270
 GCC CTC ACC ACC GCC TG 834
 Ala Leu Thr Thr Ala
 275

(2) INFORMATION FOR SEQ ID NO: 4:

(i) SEQUENCE CHARACTERISTICS

(A) LENGTH: 1209 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: double

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iii) ANTI-SENSE: NO

(iv) ORIGINAL SOURCE

(A) ORGANISM: Streptomyces

pristinaespiralis

(ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION: 1..1209

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:

ATG TCC CGT CGC CTG TTC ACC TCG GAG TCC GTG ACC GAG GGC CAC CCC	48
Met Ser Arg Arg Leu Phe Thr Ser Glu Ser Val Thr Glu Gly His Pro	
1 5 10 15	
GAC AAG ATC GCC GAC CAG ATC AGT GAC ACC GTC CTC GAC GCC CTG CTG	96
Asp Lys Ile Ala Asp Gln Ile Ser Asp Thr Val Leu Asp Ala Leu Leu	
20 25 30	
GCG GAG GAC CCC GCC TCA CGC GTC GCG GTC GAG ACC ACC CTG ATC ACC ACC	144
Arg Glu Asp Pro Ala Ser Arg Val Ala Val Glu Thr Leu Ile Thr Thr	
35 40 45	
GCG CAG GTC CAC ATC GCC GGC GAG GTC ACC ACC AAG GCG TAC GCG CCC	192
Gly Glu Val His Ile Ala Gly Glu Val Thr Thr Lys Ala Tyr Ala Pro	
50 55 60	
ATC GCC CAA CTG GTC CGC GAC ACG ATC CTG CCC ATC GGC TAC GAC TCG	240
Ile Ala Gln Leu Val Arg Asp Thr Ile Leu Ala Ile Gly Tyr Asp Ser	
65 70 75 80	
TCC GCC AAG GGC TTC GAC GGC GCC TCC TGC GGC GTC TCC GTC TCC ATC	288
Ser Ala Lys Gly Phe Asp Gly Ala Ser Cys Gly Val Ser Val Ser Ile	
85 90 95	

GCC GCG CAG TCC CCG GAC ATC GCC CAG GGC GTC GAC AGC GCC TAC GAG Gly Ala Gln Ser Pro Asp Ile Ala Gln Gly Val Asp Ser Ala Tyr Glu 100 105 110	336
ACC CGC GTC GAG GGC GAG GAC GAC GAG CTC GAC CAG CAG GGC GCC GGC Thr Arg Val Glu Gly Glu Asp Asp Glu Leu Asp Gln Gln Gly Ala Gly 115 120 125	384
GAC CAG GGC CTG ATG TTC GGC TAC GCC ACC GAC GAG ACC CCC TCG CTG Asp Gln Gly Leu Met Phe Gly Tyr Ala Thr Asp Glu Thr Pro Ser Leu 130 135 140	432
ATG CCG CTG CCC ATC GAG CTC GCC CAC CGC CTC TCG CGC CGG CTC ACC Met Pro Leu Pro Ile Glu Leu Ala His Arg Leu Ser Arg Arg Leu Thr 145 150 155 160	480
GAG GTC CGC AAG GAC GGC ACC GTC CCC TAC CTG CGC CCC GAC GGC AAG Glu Val Arg Lys Asp Gly Thr Val Pro Tyr Leu Arg Pro Asp Gly Lys 165 170 175	528
ACC CAG GTC ACC ATC GAG TAC CAG GGC AGC CGC CCG CTG CGC CTG GAC Thr Gln Val Thr Ile Glu Tyr Gln Gly Ser Arg Pro Val Arg Leu Asp 180 185 190	576
ACC GTC GTC GTC TCC TCC CAG CAC GCC GCC GAC ATC GAC CTC GGC TCC Thr Val Val Val Ser Ser Gln His Ala Ala Asp Ile Asp Leu Gly Ser 195 200 205	624
CTG CTC ACC CCC GAC ATC CGC GAG CAC GTC GTC GAG CAC GTC CTC GCC Leu Leu Thr Pro Asp Ile Arg Glu His Val Val Glu His Val Leu Ala 210 215 220	672
GCA CTC CCC GAG GAC GGC ATC AAG CTC GAG ACG GAC AAC TAC CGC CTG Ala Leu Ala Glu Asp Gly Ile Lys Leu Glu Thr Asp Asn Tyr Arg Leu 225 230 235 240	720
CTG GTC AAC CCG ACC GGC CGT TTC GAG ATC GGC GGC CCG ATG GGC GAC Leu Val Asn Pro Thr Gly Arg Phe Glu Ile Gly Gly Pro Met Gly Asp 245 250 255	768
GCC GGC CTG ACC CCC CGC AAC ATC ATC GAC ACG TAC GGC GGC ATC Ala Gly Leu Thr Gly Arg Lys Ile Ile Asp Thr Tyr Gly Gly Met 260 265 270	816
GCC CGC CAC CCC GGT GGC CGC TTC TCC GGC AAG GAC CCG TCC AAG GTC Ala Arg His Gly Gly Ala Phe Ser Gly Lys Asp Pro Ser Lys Val 275 280 285	864
GAC CGT TCC GCC GCG TAC GGC ATG CGC TGG GTC GCC AAG AAC GTC GTC Asp Arg Ser Ala Ala Tyr Ala Met Arg Trp Val Ala Lys Asn Val Val 290 295 300	912
GCC GCG GGC CTC GCC TCC CGC TGC GAG GTC CAG GTC GCC TAC GCC ATC Ala Ala Gly Leu Ala Ser Arg Cys Glu Val Gln Val Ala Tyr Ala Ile 305 310 315 320	960
GGC AAG GCC GAG CCG GTC GGC CTG TTC GTC GAG ACG TTC GGC ACC GGC Gly Lys Ala Glu Pro Val Gly Leu Phe Val Glu Thr Phe Gly Thr Gly 325 330 335	1008
ACC GTC GCC CAG GAG CGC ATC GAG AAG GCC ATC ACC GAG GTC TTC GAC Thr Val Ala Gln Glu Arg Ile Glu Lys Ala Ile Thr Glu Val Phe Asp	1056

340

345

350

CTG CGC CCC GCG GCC ATC ATC CGC GAC CTC GAC CTG CTG CGG CCC ATC 1104
Leu Arg Pro Ala Ala Ile Ile Arg Asp Leu Asp Leu Leu Arg Pro Ile
355 360 365

TAC GCC GCC ACC GCC GCC TAC GGC CAC TTC GGC CGC GAA CTG CCC GAC 1152
Tyr Ala Ala Thr Ala Ala Tyr Gly His Phe Gly Arg Glu Leu Pro Asp
370 375 380

TTC ACC TGG GAG CGG ACC GAC CGC GCC CAC CGG CTC AAG GCC GCG GCC 1200
Phe Thr Trp Glu Arg Thr Asp Arg Ala His Arg Leu Lys Ala Ala Ala
385 390 395 400

GGT CTC TG 1209
Gly Leu

(2) INFORMATION FOR SEQ ID NO: 5:

(i) SEQUENCE CHARACTERISTICS

- (A) LENGTH: 1879 base pairs
- (B) TYPE: nucleic acid
- 5 (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iii) ANTI-SENSE: NO

10 (vi) ORIGINAL SOURCE

- (A) ORGANISM: Streptomyces
pristinaespiralis

(ix) FEATURE:

(A) NAME/KEY: CDS

15 (B) LOCATION: 110..1858

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5:

GATCGGCTCC	TGACGGAGCG	GGGGCGCGCG	GGCGCGCGCG	ATCAGCGGCC	TGTCAACGGC	60
GCTGCCGACA	CTGGGGCGGA	CGCGAGGACG	AAGCCGGAAA	GGACCAACG	ATG CTG	115
					Met Leu	
					1	
GAC GGA TGC GTT CCC TGG CCC GAG GAT GTG GCC GCG AAG TAC CGG GCG						163
Asp Gly Cys Val Pro Trp Pro Glu Asp Val Ala Ala Lys Tyr Arg Ala						
5	10	15				
GCC GGC TAC TGG CGG GGC GAG CCG CTG GGC ATG CTG CTG GCC CGC TGG						211
Ala Gly Tyr Trp Arg Gly Glu Pro Leu Gly Met Leu Leu Gly Arg Trp						
20	25	30				
GCG GAG CAG TAC GGC GAG CGG GAG GCG CTG GTC GGC GCG GAC GGG TGC						259
Ala Glu Gln Tyr Gly Glu Arg Glu Ala Leu Val Gly Ala Asp Gly Cys						
35	40	45				
TCC CGT GTC ACC TAC CGT GCC CTG GAC CGC TGG TGC GAC CGG CTG GCG						307
Ser Arg Val Thr Tyr Arg Ala Leu Asp Arg Trp Cys Asp Arg Leu Ala						
55	60	65				

GCG GGG TTC GCG GCG CGC GGG ATC GGC GCC GGC GAG CCG GTG CTG GTG
 Ala Gly Phe Ala Ala Arg Gly Ile Gly Ala Gly Glu Arg Val Leu Val 355
 70 75 80

CAG CTG CCG AAC ACG CCC GAG TTC GTC GCG GTG TGC TTC GCG CTG TTC
 Gln Leu Pro Asn Thr Pro Glu Phe Val Ala Val Cys Phe Ala Leu Phe 403
 85 90 95

CGT CTG GGC GCG CTG CCG GTG TTC GCG CTG CCC GCG CAC CGT GCC GCC
 Arg Leu Gly Ala Leu Pro Val Phe Ala Leu Pro Ala His Arg Ala Ala 451
 100 105 110

GAG GTG GGG CAC CTG CTC GAG CTG TCC GGC GTC GCC CCC CAC ATC CTG
 Glu Val Gly His Leu Leu Glu Leu Ser Gly Ala Val Ala His Ile Leu 499
 115 120 125 130

CCG GGC ACC GGC ACC GGC TAC GAC CAT GTC GCG GCG GCC GTG GAG GCC
 Pro Gly Thr Gly Thr Gly Tyr Asp His Val Ala Ala Ala Val Glu Ala 547
 135 140 145

CGT GCC CGC CGT GCC CGC CCG GTG CAG GTG TTC GTG GCG GGC GAG GCG
 Arg Ala Arg Arg Ala Arg Pro Val Gln Val Phe Val Ala Gly Glu Ala 595
 150 155 160

CCC GCG GTG CTG CCC GAG GGG TTC ACC GCG CTG GCC GAC GTG GAC CCC
 Pro Ala Val Leu Pro Glu Gly Phe Thr Ala Leu Ala Asp Val Asp Gly 643
 165 170 175

GAC CCC GTG GCG CCG GCG GAC GTG GAC CCC TTC CGA CGT GCC GTC TTC
 Asp Pro Val Ala Pro Ala Asp Val Asp Ala Phe Arg Arg Gly Val Phe 691
 180 185 190

CTG CTG TCC CGG GGG ACG ACC GCG CTG CCC AAG CTG ATC CCG CGC ACC
 Leu Leu Ser Gly Gly Thr Thr Ala Leu Pro Lys Leu Ile Pro Arg Thr 739
 195 200 205 210

CAC GAC GAC TAC GCC TAC CAG TGC CGG GTC ACG GCC CGT ATC TGC CGC
 His Asp Asp Tyr Ala Tyr Gln Cys Arg Val Thr Ala Gly Ile Cys Gly 787
 215 220 225

CTG GAC GCG GAC AGT GTC TAT CTG GCG GTG CTG CCG GCC GAG TTC AAC
 Leu Asp Ala Asp Ser Val Tyr Leu Ala Val Leu Pro Ala Glu Phe Asn 835
 230 235 240

TTC CCC TTC GGC TGC CCG GGC ATC CTG GGC ACC CTG CAC GCC GGC GGG
 Phe Pro Phe Gly Cys Pro Gly Ile Leu Gly Thr Leu His Ala Gly Gly 883
 245 250 255

CGG GTG GTG TTC GCG CTG TCA CCG CAG CCC GAG GAG TGC TTC GCG CTG
 Arg Val Val Phe Ala Leu Ser Pro Gln Pro Glu Glu Cys Phe Ala Leu 931
 260 265 270

ATC GAA CGC GAA CAC GTC ACC TTC ACC TCC GTC ATC CCC ACG ATC GTG
 Ile Glu Arg Glu His Val Thr Phe Thr Ser Val Ile Pro Thr Ile Val 979
 275 280 285 290

CAC CTG TGG CTG GCG GCC GGC GCA CAA GGC CAC GGC CGC GAC CTG GGC
 His Leu Trp Leu Ala Ala Ala Gln Gly His Gly Arg Asp Leu Gly 1027
 295 300 305

AGC CTT CAG CTG CTG CAG GTC GGC AGC GCC AAA CTC CAC GAG GAG CTC
 Ser Leu Gln Leu Leu Gln Val Gly Ser Ala Lys Leu His Glu Leu 1075

183

310	315	320
GCC GCC CGG ATC GGC CCC GAA CTG GGG GTG CGG CTG CAG CAG GTG TTC Ala Ala Arg Ile Gly Pro Glu Leu Gly Val Arg Leu Gln Gln Val Phe 1123 325 . 330 335 Gly Met Ala Glu Gly Leu Leu Thr Phe Thr Arg Asp Asp Asp Pro Ala 1171 340 345 GAC GTG GTG CTG CGC ACC CAG GGC CGG CCC GTG TCC GAG GCC GAC GAG Asp Val Val Leu Arg Thr Gln Gly Arg Pro Val Ser Glu Ala Asp Glu 1219 355 360 365 ATA CGC GTC GCC GAC CCC GAC GGC CGG CCC GTG CCC CGT GAG ACC 1267 Ile Arg Val Ala Asp Pro Asp Gly Arg Pro Val Pro Arg Gly Glu Thr 375 380 385 GGT GAA CTG CTC ACC CGC GGC CCC TAC ACG CTG CCC GGC TAC TAC CGG 1315 Gly Glu Ieu Leu Thr Arg Gly Pro Tyr Thr Leu Arg Gly Tyr Tyr Arg 390 395 400 GCC CCC GAG CAC AAC GGC CGC GGC TTC ACC GAC GAC GGC TTC TAC CGC 1363 Ala Pro Glu His Asn Ala Arg Ala Phe Thr Glu Asp Gly Phe Tyr Arg 405 410 415 AGC GGC GAT CTG GTG CGG CTC ACC GGC GAC CGG CAG TTG GTG CTG GAG 1411 Ser Gly Asp Leu Val Arg Leu Thr Ala Asp Gly Gln Leu Val Val Glu 420 425 430 GGC AGG ATC AAG GAC GTC GTC ATC CGC GGC GGC GAC AAG GTC TCC CGG 1459 Gly Arg Ile Lys Asp Val Val Ile Arg Gly Asp Lys Val Ser Ala 435 440 445 450 ACC GAG GTC GAG GGC CAC CTG GGC GGC CAC CCC GAC GTC CAG CAG CGC 1507 Thr Glu Val Glu Gly His Leu Gly Ala His Pro Asp Val Gln Gln Ala 455 460 465 GCC GTC GTC GCC ATG CCC GAC CCG GTG TGG GGC GAG AAG GTC TGC GCC 1555 Ala Val Val Ala Met Pro Asp Pro Val Trp Gly Glu Lys Val Cys Ala 470 475 480 TAC ATC GTG CCC GCA CCC GGC CGT CCC GCA CCG CCG ATG GCG GCG CTG 1603 Tyr Ile Val Pro Ala Pro Gly Arg Pro Ala Pro Pro Met Ala Ala Leu 485 490 495 CGC CGG CTG CGC GCG CGG GGA CTG GGC GAC TAC AAG CTT CCC GAC 1651 Arg Arg Leu Leu Arg Ala Arg Gly Leu Ala Asp Tyr Lys Leu Pro Asp 500 505 510 CGG CTG GAG GTC GTC GAC GGC TTC CCG CTG ACC GGC CTC AAC AAG GTC 1699 Arg Val Glu Val Val Asp Ala Phe Pro Leu Thr Gly Leu Asn Lys Val 515 520 525 530 GAC AAG AAG GCC CTG GCG GCC GAC ATC GGC CCC AAG ACC GAC GGC CCC ACC 1747 Asp Lys Lys Ala Leu Ala Ala Asp Ile Ala Ala Lys Thr Ala Pro Thr 535 540 545 CGC CCC ACC ACC GCC CGC CAC GGC CGG ACC ACG GAC GGC GAT ACG GCC 1795 Arg Pro Thr Thr Ala Gly His Gly Pro Thr Thr Asp Gly Asp Thr Ala 550 555 560 GGT GGG GGT GGG TCC GCG GGC GTG ACG GGC GCG GGT GGC GGG CGG 1843 Gly Gly Gly Ser Ala Gly Gly Val Thr Ala Ala Gly Gly Gly Arg 565 570 575 GAG GAG GCG GCG TGAGCGGGCC CGGGCCCCGAG GCGCG Glu Glu Ala Ala 580 1879		

(2) INFORMATION FOR SEQ ID NO: 6:

(i) SEQUENCE CHARACTERISTICS

(A) LENGTH: 1833 base pairs
 (B) TYPE: nucleic acid
 (C) STRANDEDNESS: double
 (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iii) ANTI-SENSE: NO

(vi) ORIGINAL SOURCE

(A) ORGANISM: *Streptomyces*
pristinaespiralis

(ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION: 103..1689

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:

GGATCCCTC GCCCAGGGCC CTGGCGGCC CGCCGGGCCG TGGGGGAGGT GCGGGGGCCG 60
 CGGGCCCGG CACCGCACCA ACACAACAAC CGCTCCGGCC CC ATG CGG ACT TCA 114.
 Met Arg Thr Ser 1

CGG TCC CAC CAC CAG CGG CCC CCT ACC CCC TGG AGA CAT CCC TTG CAC 162
 Arg Ser His Asp Gln Arg Ala Pro Thr Pro Trp Arg His Pro Leu His 5 10 15 20
 5 10 15 20

AGC ACC CGG CCC GCG CCC GCC GAC CGT GAC CCC AGG CGC TGG GTC 210
 Ser Thr Arg Pro Ala Ala Asp Arg Asp Pro Arg Arg Trp Val 25 30 35
 25 30 35

ATC CTC GCC GTG ATC TGC CTG GCC CAA CTC GTC GTC CTG CTC GAC AAC 258
 Ile Leu Gly Val Ile Cys Leu Ala Gln Leu Val Val Leu Leu Asp Asn 40 45 50
 40 45 50

ACC GTC CTC AAC GTC GCC ATC CCG GTG CTC ACC ACC GAC CTG GGC GCC 306
 Thr Val Leu Asn Val Ala Ile Pro Val Leu Thr Thr Asp Leu Gly Ala 55 60 65
 55 60 65

AGC ACC GCC GAC ATC CAG TGG ATG ATC AAC GCC TAC GCG CTC GTG CAG 354
 Ser Thr Ala Asp Ile Gln Trp Met Ile Asn Ala Tyr Ala Leu Val Gln 70 75 80
 70 75 80

TCC GGG CTG CTG CTC ACC GCG GGC AGC CTC GCG GAC CGC TAC GGC CGC 402
 Ser Gly Leu Leu Leu Thr Ala Gly Ser Leu Ala Asp Arg Tyr Gly Arg 85 90 95 100
 85 90 95 100

AAA CGG CTG CTG ATG CTC GGA CTG GTG CTC TTC GGC GCC GGG TCC GCC 450

Lys Arg Leu Leu Met Leu Gly Leu Val Leu Phe Gly Ala Gly Ser Ala	105	110	115	
TGG GCG GCC TTC GCC CAG GAC TCC GCC CAA CTC ATC GCC CCC CGG GCC				498
Trp Ala Ala Phe Ala Gln Asp Ser Ala Gln Leu Ile Ala Ala Arg Ala	120	125	130	
Gly Met Gly Val Gly Ala Leu Leu Ala Thr Thr Thr Leu Ala Val	135	140	145	546
ATC ATG CAG GTC TTC GAC GAC GAA CGC CCC CGG GCG ATC GGC CTC				594
Ile Met Gln Val Phe Asp Asp Asp Glu Arg Pro Arg Ala Ile Gly Leu	150	155	160	
TGG GGA GCG GCC AGC TCA CTG GCG TTC GCG GCC CCC CGG CTC GGC				642
Trp Gly Ala Ala Ser Ser Leu Gly Phe Ala Ala Gly Pro Leu Leu Gly	165	170	175	
180				
GCC GCC CTC CTC GAC CAC TTC TGG TGG CCC TCC ATC TTC CTG ATC AAC				690
Gly Ala Leu Leu Asp His Phe Trp Trp Gly Ser Ile Phe Leu Ile Asn	185	190	195	
CTG CCC GTC GCG CTG CTG GGC CTG CTG GCC GTC GCC CCC CTG GTG CCC				738
Leu Pro Val Ala Leu Leu Gly Leu Ala Val Ala Arg Leu Val Pro	200	205	210	
GAG ACG AAG AAC CCC GAA GCG CGG CCC CCC GAC CTG CTC GGC CCC CTG				786
Glu Thr Lys Asn Pro Glu Gly Arg Arg Pro Asp Leu Leu Gly Ala Val	215	220	225	
CTC TCC ACC CTC GCC ATG GTC GGC GTC GTC TAC GCC ATC ATC TCC GCC				834
Leu Ser Thr Leu Gly Met Val Gly Val Val Tyr Ala Ile Ile Ser Gly	230	235	240	
CCC GAA CAC CCC TGG ACG GCC CCG CAG GTC CTC CTG CCC CCC GGC GTC				882
Pro Glu His Gly Trp Thr Ala Pro Gln Val Leu Leu Pro Ala Ala Val	245	250	255	
260				
GCG GCC GCG CTC ACC CCG TTC GTC CGC TGG GAA CTG CAC ACC CCC				930
Ala Ala Ala Leu Thr Ala Phe Val Arg Trp Glu Leu His Thr Pro	265	270	275	
CAC CCC ATG CTC GAC ATG GCC TTC TTC ACC GAC CGG CGC TTC AAC GGG				978
His Pro Met Leu Asp Met Gly Phe Phe Thr Asp Arg Arg Phe Asn Gly	280	285	290	
CCG TCG CCG GCG GAG TGC TCG TCG TTC GGC ATG GCC GGC TCG CTC TTC				1026
Pro Ser Pro Ala Glu Cys Ser Ser Phe Gly Met Ala Gly Ser Leu Phe	295	300	305	
CTG CTC ACC CAG CAC CTC CAA CTC GTC CTC GGC TAC GAC GCC CTG CAG				1074
Leu Leu Thr Gln His Leu Gln Leu Val Leu Gly Tyr Asp Ala Leu Gln	310	315	320	
GCC GGC CTG CGC ACC GCG CCA CTG GCT TTG ACG ATC GTC GCC CTC AAC				1122
Ala Gly Leu Arg Thr Ala Pro Leu Ala Leu Thr Ile Val Ala Leu Asn	325	330	335	
340				
CTG GCC GGC CTC GGC GCG AAA CTC CTC GCC GCG CTC GGC ACC GCC CGC				1170
Leu Ala Gly Leu Gly Ala Lys Leu Leu Ala Ala Leu Gly Thr Ala Arg	345	350	355	

AGC ATC GCC CTG GGC ATG ACA CTG CTC GCC GGC CTC AGC GCG GTG 1218
 Ser Ile Ala Leu Gly Met Thr Leu Leu Ala Ala Gly Leu Ser Ala Val
 360 365 370
 GCC GTC GGC GGA TCG GGC CCC GAC GCC GGC TAC GGC GGC ATG CTC GCC 1266
 Ala Val Gly Gly Ser Gly Pro Asp Ala Gly Tyr Gly Gly Met Leu Ala
 375 380 385
 GCC CTG CTC CTC ATG GGC GCG GGC ATC GCA CTG GCC ATG CCC GCC ATG 1314
 Gly Leu Leu Met Gly Ala Gly Ile Ala Leu Ala Met Pro Ala Met
 390 395 400
 GCC ACC GCC GTG ATG TCC TCC ATC CCG CCC GCC AAG GCC GGG GCC GGA 1362
 Ala Thr Ala Val Met Ser Ser Ile Pro Pro Ala Lys Ala Gly Ala Gly
 405 410 415 420
 GCC GGC GTG CAG GGC ACC CTG ACC GAG TTC GGC GGC GGA CTG GGA GTG 1410
 Ala Gly Val Gln Gly Thr Leu Thr Glu Phe Gly Gly Leu Gly Val
 425 430 435
 GCC ATC CTC GCC GGC CTC CTC GCC TCC CCC TTC GGC TCC CAA CTG CCC 1458
 Ala Ile Leu Gly Ala Val Leu Gly Ser Arg Phe Ala Ser Gln Leu Pro
 440 445 450
 GCC GCC ATC ACC ACC GGC ACC TCC CTC GAC GAG GCA CTG CCC GAC GGC 1506
 Ala Ala Ile Thr Gly Thr Gly Ser Leu Asp Glu Ala Leu Arg Asp Ala
 455 460 465
 ACA CCC CAA CAG GCC GGG CAG GTC CAC GAC GCG TTC GCC GAC GCG GTG 1554
 Thr Pro Gln Gln Ala Gly Gln Val His Asp Ala Phe Ala Asp Ala Val
 470 475 480
 AAC ACC AGC CAA CTC ATC GGC GCC CCC GGC GTG TTC ACC GGC GGC CTG 1602
 Asn Thr Ser Gln Leu Ile Gly Ala Ala Ala Val Phe Thr Gly Leu
 485 490 495 500
 CTC GCC GCG CTG CTG CTC CAC CGC GCC GAC CGC AAG GCC GCC CCC CAG 1650
 Leu Ala Ala Leu Leu Leu His Arg Ala Asp Arg Lys Ala Ala Pro Gln
 505 510 515
 CCC ACC GCC CCC ACC CCC GAA CCC ACC ACC ACC GCC TGACCCCCGG 1696
 Pro Thr Ala Pro Thr Pro Glu Pro Thr Thr Thr Ala
 520 525
 CCCGGCGGGC ACCACACAAAC CCACGGCCCC ACCCTGCGG CTCCCCACCG GGACCCACAG 1756
 GGCGGGGGCC GTGCCGCTGC CCTGCCACA CACACAGCCC CCACACACAC AGCCCCGGCA 1816
 CGGCCGACAG CGCCGGGG
 1833

(2) INFORMATION FOR SEQ ID NO: 7:

(i) SEQUENCE CHARACTERISTICS

(A) LENGTH: 695 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: double

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iii) ANTI-SENSE: NO

(vi) ORIGINAL SOURCE

5 (A) ORGANISM: *S. pristinaespiralis*

(ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION: 212..695

(D) OTHER INFORMATION: /product = "SnaC Gene"

10 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7:

CTCGAGCCCCC	GCCCCCAGGT	CCTGGTGTG	CTCGCCGTGG	AGAAGGGCGC	CGACGGCACC	60					
GCGCCGCCGG	ACCGGCTGCT	GATCCACGAC	GGCTTCCCCCT	GGGGCCGCGC	CGCCCCCGCGC	120					
GAAGCGGAGC	TGCCCAACCGG	GCACCCGCGCC	CTGCCGGCCC	TGGCCGGCGC	CGCCCGCTGA	180					
GGCGCGGCAA	CCACCAACAG	AAGGAGCCCC	C	GTG	ACA	GGA	CCC	GAC	GAC	CCG	232
				Val	Thr	Gly	Ala	Asp	Asp	Pro	
				1				5			

15 GCA AGG CCC GCG GTC GGC CCG CAG ACT TTC CGA GAC GCG ATG GCG CAG 280
Ala Arg Pro Ala Val Gly Pro Gln Ser Phe Arg Asp Ala Met Ala Gln
10 15 20

CTG GCG TCG CCC GTC ACC GTC GTA ACC GTC CTC GAC GCG GCC CGA CGC 328
 Leu Ala Ser Pro Val Thr Val Val Thr Val Leu Asp Ala Ala Gly Arg
 25 30 35

CGC CAC GGC TTC ACG GCC GGC TCG GTG GTC TCT GTG TCG CTG GAC CCG 376
 Arg His Gly Phe Thr Ala Gly Ser Val Val Ser Val Ser Leu Asp Pro
 40 45 50 55

CCG CTG GTG ATG GTC GGC ATC GCG CTC ACC TCC AGC TGC CAC ACG GCG 424
 Pro Leu Val Met Val Gly Ile Ala Leu Thr Ser Ser Cys His Thr Ala
 60 65 70

ATG GCC GCC GCC GAG TTC TGC GTC AGC ATC CTC GGC GAG GAC CAG 472
Met Ala Ala Ala Ala Glu Phe Cys Val Ser Ile Leu Gly Glu Asp Gln
75 80 85

Arg Ala Val Ala Lys Arg Cys Ala Thr His Gly Ala Asp Arg Phe Ala
90 95 100

Gly Gly Glu Phe Ala Ala Trp Asp Gly Thr Gly Val Pro Tyr Leu Pro
105 110 115 . 568

GAC GCC AAG GTC GTC CGG CGC TGC CGC ACC ACG GAC GTC GTG CGC GCC 616
 Asp Ala Lys Val Val Leu Arg Cys Arg Thr Thr Asp Val Val Arg Ala
 120 125 130 135

GGC GAC CAC GAC CTG GTG CTC GGC ACG CCC GTG GAG ATC CGC ACG GGC
 Gly Asp His Asp Leu Val Leu Gly Thr Pro Val Glu Ile Arg Thr Gly
 140 145 150
 GAC GGC GCG AAC GCA CGG GTC GCG GTC G
 151

Asp Pro Ala Lys Pro Pro Leu Leu Trp Tyr
155 160

(2) INFORMATION FOR SEQ ID NO: 8:

(i) SEQUENCE CHARACTERISTICS

(A) LENGTH: 640 base pairs

(B) TYPE: nucleic acid

5 (C) STRANDEDNESS: double

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iii) ANTI-SENSE: NO

10 (vi) ORIGINAL SOURCE

(A) ORGANISM: *S. pristinaespiralis*

(ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION: 1..640

15 (D) OTHER INFORMATION: /product = "SnaD gene"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 8:

GGG ACC GCC CGG CTC ATC GGC CCG CTG CCG CGG CTG GGC CTC CAG	48
Ala Thr Ala Arg Leu Ile Gly Pro Leu Pro Arg Arg Leu Gly Leu Gln	
1 5 10 15	

GTC CAC CAG GTG ATG ACG GGC GCG TTC GCG CAG GCC CTC GCC CGC TGG	96
Val His Gln Val Met Thr Gly Ala Phe Ala Gln Ala Leu Ala Arg Trp	
20 25 30	

CGG GGC AGC CGC GCC GTC ACC TTC GAC GTG GAG ACC CAC GGA CGG CAC	144
Arg Gly Ser Arg Ala Val Thr Phe Asp Val Glu Thr His Gly Arg His	
35 40 45	

CCC CGC CAC GAA CTG TTC CGT ACC GTC GGC TGG TTC ACC TCC ATC CAC	192
Gly Arg Asp Glu Leu Phe Arg Thr Val Gly Trp Phe Thr Ser Ile His	
50 55 60	

CCC GTC GTC CTG CGC GCG GAC CGC TCC GTG CAC CCC GAG CAG TAC CTC	240
Pro Val Val Leu Gly Ala Asp Arg Ser Val His Pro Glu Gln Tyr Leu	
65 70 75 80	

GCC CAG ATC GGC GCG GCG CTG ACC GCC GTC GAA CCG GAC GGC GGC GTC CGC	288
Ala Gln Ile Gly Ala Ala Leu Thr Ala Val Pro Asp Gly Gly Val Gly	
85 90 95	

TTC CGC GCC TGC CGC GAG TTC TCC CCG GAC GCC GGG CTG CGC ACT CTC	336
Phe Gly Ala Cys Arg Glu Phe Ser Pro Asp Ala Gly Leu Arg Thr Leu	
100 105 110	

CTG CGT GAC CTG CCG CCC CCC CTG GTG TGC TTC AAC TAC TAC GGT CAG	384
Leu Arg Asp Leu Pro Pro Ala Leu Val Cys Phe Asn Tyr Tyr Gly Gln	
115 120 125	

GCC GAC CAG TTG AGC CCG AAC GGC GGT TTC CGT ATG TCG GGC CGT CCC Ala Asp Gln Leu Ser Pro Asn Gly Gly Phe Arg Met Ser Gly Arg Pro 130 135 140	432
ATC CCG CGC GAG CAC TCC GCC CGC TGC GAG CGC GTC TAC GGC ATC GAG Ile Pro Arg Glu His Ser Ala Arg Cys Glu Arg Val Tyr Gly Ile Glu 145 150 155 160	480
GTC TAC GGC ATC GTC CAC GGC CGC CTG CGC ATG GGC CTG ACC TGG Val Tyr Gly Ile Val His Gly Arg Leu Arg Met Gly Leu Thr Trp 165 170 175	528
GTC CGG AGC CCG GCG GAC GGT GTG GAC GAG GCC GGC GTC GAC GCG CTC Val Ile Ser Pro Ala Asp Gly Val Asp Glu Ala Gly Val Asp Ala Leu 180 185 190	576
GTC GAG CAG ATG AGC TGG GTG CTG GCC ACG CTC GCG GGC GCC GAC CCG Val Glu Gln Met Ser Trp Val Leu Ala Thr Leu Ala Gly Ala Asp Pro 195 200 205	624
CAC GCC GTG ACC CCG G His Ala Val Thr Pro 210	640

(2) INFORMATION FOR SEQ ID NO: 9:

(i) SEQUENCE CHARACTERISTICS

(A) LENGTH: 645 base pairs

(B) TYPE: nucleic acid

5 (C) STRANDEDNESS: double

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iii) ANTI-SENSE: NO

10 (vi) ORIGINAL SOURCE

(A) ORGANISM: *S. pristinaespiralis*

(ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION: 61..645

15 (D) OTHER INFORMATION: /product = "papA gene"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:

190

GCGGTCAAGA ACCTGCCGCT GACCGTACGG CGCCGCTGAC ACAGACAAGG GGCCACCTG 60
 GTG CGC ACC GTG CGA ACC CTG CTG ATC GAC AAC TAC GAC TCG TTC ACC
 Val Arg Thr Val Arg Thr Leu Leu Ile Asp Asn Tyr Asp Ser Phe Thr 108
 1 5 10 15
 TAC AAC CTC TTC CAG ATG CTG GCC GAG GTG AAC GCC CCC GCT CCG CTC 156
 TYR Asn Leu Phe Gln Met Leu Ala Glu Val Asn Gly Ala Ala Pro Leu
 20 25 30
 GTC GTC CGC AAC GAC GAC ACC CCC TGG CAG CCC CTG GCG CCG CCC 204
 Val Val Arg Asn Asp Asp Thr Arg Thr Trp Gln Ala Leu Ala Pro Gly
 35 40 45
 GAC TTC GAC AAC GTC GTC TCA CCC GGC CCC GGC CAC CCC CCC ACC 252
 Asp Phe Asp Asn Val Val Val Ser Pro Gly Pro Gly His Pro Ala Thr
 50 55 60
 GAC ACC GAC CTG GGC CTC AGC CGC CGG GTG ATC ACC GAA TGG GAC CTG 300
 Asp Thr Asp Leu Gly Leu Ser Arg Arg Val Ile Thr Glu Trp Asp Leu
 65 70 75 80
 CCG CTG CTC GGG GTG TGC CTG GGC CAC CAG CCC CTG TGC CTG CTC CCC 348
 Pro Leu Leu Gly Val Cys Leu Gly His Gln Ala Leu Cys Leu Ala
 85 90 95
 GCC GCC GCC GTC GTC CAC GCA CCC GAA CCC TTT CAC GGC CGC ACC AGC. 396
 Gly Ala Ala Val Val His Ala Pro Glu Pro Phe His Gly Arg Thr Ser
 100 105 110
 GAC ATC CCC CAC GAC GGG CAG GGC CTG TTC GGC AAC ATC CCC TCC CCC 444
 Asp Ile Arg His Asp Gly Gln Gly Leu Phe Ala Asn Ile Pro Ser Pro
 115 120 125
 CTG ACC GTG GTC CGC TAC CAC TCG CTG ACC GTC CGG CAA CTG CCC CCC 492
 Leu Thr Val Val Arg Tyr His Ser Leu Thr Val Arg Gln Leu Pro Ala
 130 135 140
 GAC CTG CGC CCC ACC GCC CAC ACC GCC GAC GGG CAG CTG ATG GCC GTC 540
 Asp Leu Arg Ala Thr Ala His Thr Ala Asp Gly Gln Leu Met Ala Val
 145 150 155 160
 GCC CAC CGC CAC CTG CCC CCC TTC GGC GTG CAG TTC CAC CCC GAA TCG 588
 Ala His Arg His Leu Pro Arg Phe Gly Val Gln Phe His Pro Glu Ser
 165 170 175
 ATC ACC ACC GAA CAC GGC CAC CGG ATG CTC CCC AAC TTC CGC GAC CTG 636
 Ile Ser Ser Glu His Gly His Arg Met Leu Ala Asn Phe Arg Asp Leu
 180 185 190
 TCC CTG CGC
 Ser Leu Arg
 195

(2) INFORMATION FOR SEQ ID NO: 10:

(i) SEQUENCE CHARACTERISTICS

- (A) LENGTH: 1052 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iii) ANTI-SENSE: NO

(vi) ORIGINAL SOURCE

5

(A) ORGANISM: *S. pristinaespiralis*

(ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION: 84-962

(D) OTHER INFORMATION: /product = "pamM gene"

10

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 10:

CTCAGGAGCC ACTGGATCCC CTCCCCCCCCC GCCCCCTCC CCACCCCCCTTCC 60
 GCGTCCCCCGC GGGGGGCGGT GTC GTC ACT CCC CCC GCA CCC ACC CTC CCC 110
 Val Thr Ala Ala Ala Pro Thr Leu Ala

GGC GAC GCG CCC CGG GCG GAC ACC CGG CTG CTG GCG GCG GAC GCG TGC Ala Asp Ala Ala Arg Ala Asp Thr Arg Leu Leu Ala Ala His Ala Cys	206	
30	35	40
CAG GTC GCG CCC CGG GCG GAC GTC GAC ACC TGC CGG CCC GCG GCG CTG GCG Gln Val Ala Pro Gly Asp Leu Asp Thr Cys Leu Ala Gly Pro Val Pro	254	
45	50	

CCC CGG TTC TCG CAC TAC GTC CGG CCC CGT CTG ACC CCC GAA CCC CCC
 Pro Arg Phe Tyr His Tyr Val Arg Arg Arg Leu Thr Arg Glu Pro Ala
 60 65 70

GAA CGC ATC GTC GGC CAC CCC TAC TTC ATG GGC CAC CGC TTC GAC CTG
 Glu Arg Ile Val Gly His Ala Tyr Phe Met Gly His Arg Phe Asp Leu.
 75 80 85

GGT CCC GGC GTC TTC GTC CCC AAA CCC GAG ACC GAG GAG ATC ACC CCC	398
Ala Pro Gly Val Phe Val Pro Lys Pro Glu Thr Glu Glu Ile Thr Arg	
90 95 100 105	
GAC GGC ATC GGC CCC CTC GAG GGC CTC CTC CCC CCC GGC ACC ACC CCA	446
Asp Ala Ile Ala Arg Leu Glu Ala Leu Val Arg Arg Gly Thr Thr Ala	
110 115 120	

CGC CTC GTC GTC GAC CTC TCC CCC GAA CCG CGC ACC ATC GCG CTC ACC 494
 PRO Leu Val Val Asp Leu Cys Ala Gly Pro Gly Thr Met Ala Val Thr
 125 130 135
 CTG CGC CGC CAC GIA CGC CGC CGC GTC CTC CGC CGC ATC GAA CTC TCC 542
 Leu Ala Arg His Val Pro Ala Ala Arg Val Leu Gly Ile Glu Leu Ser
 140 145

CAG CCC CCC CCC CCC CCC CCC AAC CCC CCC CCC ACC CCC CCC 590
 Gln Ala Ala Ala Arg Ala Ala Arg Arg Asn Ala Arg Gly Thr Gly Ala
 155 160 165

CCG ATC CTC CAC CCC CCC GAC CCC TTC CCC GAA CTG ACC CGC	638
Arg Ile Val Gln Gly Asp Ala Arg Asp Ala Phe Pro Glu Leu Ser Gly	
170 175 180 185	
ACC CTC CAC CTC CTC ACC AAC CCC CCC TAC ATC CCC ATC CGA CTG	686
Thr Val Asp Leu Val Val Thr Asn Pro Pro Tyr Ile Pro Ile Gly Leu	

190	195	200
CCC ACC TCC GCA CCC GAA CTC CTC GAG CAC GAC CCC CCC CTC CCC CTC		
Arg Thr Ser Ala Pro Glu Val Leu Glu His Asp Pro Pro Leu Ala Leu		
205	210	215

TGG GCC GGG GAG GAC GGC CTC GGC ATG ATC CGC GCC ATG GAA CGC ACC 782
 Trp Ala Gly Glu Glu Gly Leu Gly Met Ile Arg Ala Met Glu Arg Thr

220 225 230

GCG GCC CGG CTG CTG GCC CCC GGC GGC GTC CTG CTC CTC GAA CAC GGC 830
 Ala Ala Arg Leu Leu Ala Pro Gly Gly Val Leu Leu Leu Glu His Gly
 235 240 245

TCC TAC CAA CTC GCC TCC GTG CCC GCC CTG TTC CGC GCA ACC GGC CGC 878
 Ser Tyr Gln Leu Ala Ser Val Pro Ala Leu Phe Arg Ala Thr Gly Arg
 250 255 260 265

TGG AGC CAC CCC TCG TCC CGT CCC ACC TGC AAC GAC GGC TGC CTG ACC 926
 Trp Ser His Ala Ser Ser Arg Pro Thr Cys Asn Asp Gly Cys Leu Thr
 270 275 280

GCC GTC CGC AAC CAC ACC TGC GCA CCG CCC TGACACGGCG TCACGGCAGG 979
 Ala Val Arg Asn His Thr Cys Ala Pro Pro Ala
 285 290

GCGGGCTGT CGGCAACGAC CCTACCCAT TGACAAACCG ACCGTGCCGT TTTTTAATG 1039
 TCGGGGTGGC GGA
 1052

(2) INFORMATION FOR SEQ ID NO: 11:

(i) SEQUENCE CHARACTERISTICS

(A) LENGTH: 227 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: double

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO

(vi) ORIGINAL SOURCE

(A) ORGANISM: *S. pristinaespiralis*

(ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION: 3..227

(D) OTHER INFORMATION: /product = "Part of
 the SnbC gene"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 11:

AG ATC TTC GAG CAC AAG ACC GTC GCC CAG CTC GCA CCC GTC GCC GAG Ile Phe Glu His Lys Thr Val Ala Gln Leu Ala Pro Val Ala Glu	47
1 5 10 15	
ACG CTC GCC GAC ACC ACC CGC GAG GAA CCC GCC GTC GCC GCG ACC Thr Leu Ala Asp Thr Thr Arg Glu Glu Pro Ala Ala Val Ala Ala Thr	95
20 25 30	
GGC GAC GTA CCG CTC ACC CCG ATC ATG CAC TGG CTG CGC GAA CGC GGC Gly Asp Val Pro Leu Thr Pro Ile Met His Trp Leu Arg Glu Arg Gly	143
35 40 45	
GGC CCC GTC GAC GCG TTC AGC CAG ACC ATG CCC GTC ACC GTC CCC GCG Gly Pro Val Asp Ala Phe Ser Gln Thr Met Ala Val Thr Val Pro Ala	191
50 55 60	
GGC CTG GAC CGG GAA CGG CTC GTG GCC CCC CTG CAG Gly Leu Asp Arg Glu Arg Leu Val Ala Ala Leu Gln	227
65 70 75	

(2) INFORMATION FOR SEQ ID NO: 12:

(i) SEQUENCE CHARACTERISTICS

(A) LENGTH: 247 base pairs

(B) TYPE: nucleic acid

5 (C) STRANDEDNESS: double

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iii) ANTI-SENSE: NO

10 (vi) ORIGINAL SOURCE

(A) ORGANISM: *S. pristinaespiralis*

(ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION: 1..247

(D) OTHER INFORMATION: /product = "Part of the SnbC gene"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 12:

CTC GAG TAC GAC ACC GCC CTG TAC GAG CGG GCC ACC GCC GAA GCC CTC	48
Leu Glu Tyr Asp Thr Ala Leu Tyr Glu Arg Ala Thr Ala Glu Ala Leu	
1 5 10 15	
ACC GGC CGG CTG CTG CGG CTC CTC GAC GCC GTC GTC ACC GAC CCG CAG	96
Thr Gly Arg Leu Leu Arg Leu Leu Asp Ala Val Val Thr Asp Pro Gln	
20 25 30	
GCG CCG GTC GGC TCC CAC GAC CTC CTC GAA GAG GCC GAA CAC GCC CGC	144
Ala Pro Val Gly Ser His Asp Leu Leu Glu Ala Glu His Ala Arg	
35 40 45	
CTG GCA GCC TTC AAC GAC ACC GCC CGG CCC GTG CCG CGA GCC GGC CTC	192
Leu Ala Ala Phe Asn Asp Thr Ala Arg Pro Val Pro Arg Ala Gly Leu	
50 55 60	
GCC GAA CTC TTC ACC CCC CAG GCC CCC CGC ACC GCC GAT GCG GTC GCC	240
Ala Glu Leu Phe Thr Ala Gln Ala Arg Arg Thr Ala Asp Ala Val Ala	
65 70 75 80	
GTC GTC G	247
Val Val	

5 (2) INFORMATION FOR SEQ ID NO: 13:

(i) SEQUENCE CHARACTERISTICS

(A) LENGTH: 192 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: double

10 (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iii) ANTI-SENSE: NO

(vi) ORIGINAL SOURCE

15 (A) ORGANISM: *S. pristinaespiralis*

(ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION: 3..192

5

(D) OTHER INFORMATION: /product = "Part of
the SnbD gene"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 13:

GC ATG CCC CCC GTC ACC CCC TAC CGC GCC TAC CTG GCC CAC CTC CCC Met Pro Pro Val Thr Pro Tyr Arg Ala Tyr Leu Ala His Leu Ala	47
1 5 10 15	
GGC CGT GAC GAC GAC GCC CGC GCC GCG TGG CGG ACC GCC CTC CGC Gly Arg Asp Asp Asp Ala Ala Arg Ala Ala Thr Arg Thr Ala Leu Ala	95
20 25 30	
GAC CTG GAG GAG CCG AGC CTC GTC GCG GGC GCC GGA GCA GGC CGC CCC Asp Leu Glu Glu Pro Ser Leu Val Ala Gly Ala Gly Ala Gly Arg Gly	143
35 40 45	
GCC GCC GAC CCC TCC CCC CTG CCC GGC CAG ATC CCC GGT TAC CGA GCT C Ala Ala Asp Gly Ser Ala Leu Pro Gly Gln Ile Pro Gly Tyr Arg Ala	192
50 55 60	

(2) INFORMATION FOR SEQ ID NO: 14:

(i) SEQUENCE CHARACTERISTICS

(A) LENGTH: 474 base pairs

10

(B) TYPE: nucleic acid

(C) STRANDEDNESS: double

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

15

(iii) ANTI-SENSE: NO

(vi) ORIGINAL SOURCE

(A) ORGANISM: S. pristinaespiralis

(ix) FEATURE:

(A) NAME/KEY: CDS

5

(B) LOCATION: 1..474

(D) OTHER INFORMATION: /product = "Part of
the SnbD gene"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 14:

CTG CAG GTC GAG GGC CGG CCC GCG CAC CTG GAA CTG CCC TGC GAC CAC	48
Leu Gln Val Glu Gly Arg Pro Ala His Leu Glu Leu Pro Cys Asp His	
1 5 10 15	
CCC CGG CCC GCC GTC GCC ACC CAC CCC GGC GCC ACC GTG CCC TTC CAC	96
Pro Arg Pro Ala Val Ala Thr His Arg Gly Ala Thr Val Pro Phe His	
20 25 30	
ATC GAC GCC GCC CTC CAC CAG AAG CTG ACC GGC CTC TCC AAG GCA TGC	144
Ile Asp Ala Gly Leu His Glu Lys Leu Thr Ala Leu Ser Lys Ala Cys	
35 40 45	
GAC AGC AGC CTG TTC ATG GTG CTC CAG GCC GGC GTC GCC CCC CTG CTC	192
Asp Ser Ser Leu Phe Met Val Leu Gln Ala Ala Val Ala Ala Leu Leu	
50 55 60	
ACC CGG CAC GGC GCC GGC ACC GAC ATC CCC GTC GGC AGC CCC GTC GCC	240
Thr Arg His Gly Ala Gly Thr Asp Ile Pro Val Gly Ser Pro Val Ala	
65 70 75 80	
GGC CGC ACC GAC GAC GCC CTC GAC GAC CTG CTG GGC TTC TTC GTC AAC	288
Gly Arg Thr Asp Asp Ala Leu Asp Asp Leu Val Gly Phe Phe Val Asn	
85 90 95	
ACC CTC GTC CTG CGC ACC GAC ACC TCC GGC GAC CCC ACC TTC CGC GAA	336
Thr Leu Val Leu Arg Thr Asp Thr Ser Gly Asp Pro Thr Phe Arg Glu	
100 105 110	
CTC GTC GCA CGC GTG CGG CAG TTC GAC CTC CCC GGC TAC ACG CAC CAG	384
Leu Val Ala Arg Val Arg Gln Phe Asp Leu Ala Ala Tyr Thr His Gln	
115 120 125	
GAC ATG CCG-TTC GAA AAG CTC GTC GAA CAG GTC AAC CCC GAC CGC TCC	432
Asp Met Pro Phe Glu Lys Leu Val Glu Glu Val Asn Pro Glu Arg Ser	
130 135 140	
CTG GCC CGC AAC CCG CTC TTC CAG GTC GTC CTG GCG CTG CAG	474
Leu Ala Arg Asn Pro Leu Phe Gln Val Val Leu Ala Leu Gln	
145 150 155	

(2) INFORMATION FOR SEQ ID NO: 15:

(i) SEQUENCE CHARACTERISTICS

(A) LENGTH: 485 base pairs

(B) TYPE: nucleic acid

5 (C) STRANDEDNESS: double

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iii) ANTI-SENSE: NO

10 (vi) ORIGINAL SOURCE

(A) ORGANISM: *S. pristinaespiralis*

(ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION: 3..485

15 (D) OTHER INFORMATION: /product = "Part of
the SnbE gene"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 15:

GC ATG CCG CGC TCC CTC GAC CTG TAC GTC GCA CTG CTC GCC GTC CTC Met Pro Arg Ser Leu Asp Leu Tyr Val Ala Leu Leu Ala Val Leu	47
1 5 10 15	
AAG ACC GGC GCC GCC TAC CTG CCC GTC GAC ATC TCC TAC CCG GCC GAA Lys Thr Gly Ala Ala Tyr Leu Pro Val Asp Ile Ser Tyr Pro Ala Glu	95
20 25 30	
CGC ATC CGG TTC ATG ATC GAG GAC GCC CGC CCG GTG ACC GTC CTC GAC Arg Ile Ala Phe Met Ile Glu Asp Ala Arg Pro Val Thr Val Leu Asp	143
35 40 45	
CGC CTG CCC GAC GAC CTG GGC GCC TAC CGG GAC ACC GAC CTC ACC GAC Arg Leu Pro Asp Asp Leu Gly Ala Tyr Arg Asp Thr Asp Leu Thr Asp	191
50 55 60	
GCC GAC CGC ACG GCG CCG CTA CGG CCC GAA CAC CCG GCG TAC GTC ATC Ala Asp Arg Thr Ala Pro Leu Arg Pro Glu His Pro Ala Tyr Val Ile	239
65 70 75	
CAC ACC TCC GGC TCC ACC GGC ACC CCC AAG GGC GTC GTC ATG CCC CAC His Thr Ser Gly Ser Thr Gly Thr Pro Lys Ala Val Val Met Pro His	287
80 85 90 95	
GCC GGC CTG GTC AAC CTG CTG ACC TGG CAC GCC CGC CGC TTC CCC GGC Ala Gly Leu Val Asn Leu Leu Thr Trp His Ala Arg Arg Phe Pro Gly	335
100 105 110	

GGC ACC GGG GTG CGC ACC GCC CAG TTC ACC GCC ATC GGC TTC GAC TTC	383
Gly Thr Gly Val Arg Thr Ala Gln Phe Thr Ala Ile Gly Phe Asp Phe	
115 120 125	
TCG GTG CAG GAG ATC CTC TCC CCG CTC GTC ATG GGC AAG ACC CTC GCC	431
Ser Val Gln Glu Ile Leu Ser Pro Leu Val Met Gly Lys Thr Leu Ala	
130 135 140	
TG C CCC TCG GAA GAG GTC CGC CAC ACC CCC GAA CTG CTG CCC CCC TGG	479
Val Pro Ser Glu Glu Val Arg His Ser Ala Glu Leu Leu Ala Gly Trp	
145 150 155	
CTC GAG	485
Leu Glu	
160	

(2) INFORMATION FOR SEQ ID NO: 16:

(i) SEQUENCE CHARACTERISTICS

(A) LENGTH: 291 base pairs

(B) TYPE: nucleic acid

5 (C) STRANDEDNESS: double

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO

10 (vi) ORIGINAL SOURCE

(A) ORGANISM: *S. pristinaespiralis*

(ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION: 1..291

15 (D) OTHER INFORMATION: /product = "Part of
the SnbE gene"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 16:

CTG CAG GCC GAC CCC GCC GAA GTG AGC CTG CTG GCC GTC CTC GAC GCC	48
Leu Gln Ala Glu Gly Ala Glu-Val Ser Leu Leu Ala Val Leu Asp Gly	
1 5 10 15	
TAC CCC GAC GCC TAC GAC CCC ACC GAG CAC GAG GTC GGC GAG GAA CAG	96
Tyr Pro Asp Ala Tyr Asp Gly Thr Glu His Glu Val Gly Glu Glu Gln	
20 25 30	
GTC CTG GCG ATC CTC CTC AAC GCC CCC GGC GTC GAC CGG GCC CAG CCC	144
Val Leu Ala Ile Leu Leu Asn Ala Ala Gly Val Asp Arg Ala Gln Ala	
35 40 45	
TTC GGC GAC CCC CTC CAA CGG GCC GCC GTG CTC GAG AAG CTG CGC	192
Phe Gly Asp Ala Pro Leu Gln Arg Ala Ala Val Leu Glu Lys Leu Arg	
50 55 60	
GAC AGC GGC AGC GCC CTG GGC AAC CTC GAC GAC GCG GTC GGC CGC	240
Asp Ser Gly Ser Ala Leu Gly Asn Leu Asp Asp Asp Ala Val Gly Arg	
65 70 75 80	
ATG GTC ACC GTC TTC CTC AAC AAC ACG CCC CTC ATC CAG AAC TTC CGG	288
Met Val Thr Val Phe Leu Asn Asn Thr Arg Leu Ile Gln Asn Phe Arg	
85 90 95	
CCC	291
Pro	